	Ш
	Ш
	Ш
	齫
	W
在一个人的主义,只要是一个人的主义,但是一个人的主义,但是一个人的主义,但是一个人的主义,但是一个人的主义,但是一个人的主义,但是一个人的主义,但是一个人的主义 第二章 1955年,1955年,1955年,1955年,1955年,1955年,1955年,1955年,1955年,1955年,1955年,1955年,1955年	SHEET

In sife III

RE 1 N365 1986

ANNUAL REPORT
National Eye Institute
October 1, 1985 - September 30, 1986

REPORT OF THE SCIENTIFIC DIRECTOR Jin H. Kinoshita, Ph.D.

During this past year considerable restructuring of the intramural program has taken place. The Laboratory of Vision Research (LVR) has been divided into smaller working units. During the course of years the LVR accumulated a number of unrelated research groups making the Laboratory difficult to manage. Creating smaller units of scientists with similar interests strengthens the interaction and helps the head of the Laboratory to be more responsive to the needs of his group. Three additional laboratories, Laboratory of Retinal Cell and Molecular Biology, Laboratory of Immunology and Laboratory of Mechanisms of Ocular Diseases have been created. These laboratories join the Clinical Branch, Laboratory of Sensorimotor Research, Laboratory of Molecular and Developmental Biology and Laboratory of Pathology to constitute the seven branches of the NEI Intramural Program. We feel the reorganization makes for a more efficient means of managing the research activities on the Bethesda campus.

During the year significant progress has been achieved in many research areas. To illustrate this point four studies will be cited in this report even though there are several other intramural studies which are equally impressive.

For a number of years Dr. Robert Nussenblatt, Dr. Igal Gery and their associates have been developing experimental animals models which simulate the inflammatory eye diseases known as uveitis. The most promising approach was found to be the injection of a purified retinal protein, identified as S antigen, into Lewis rats, guinea pigs and monkeys. Animals immunized with S antigen developed clinical anterior and posterior uveitis which was confirmed by histology. monkey model closely resembled the disease found in posterior uveitis patients. Additional experimental study suggested to the group that T-cells were involved in this ocular inflammatory process. Since cyclosporine, a complex peptide isolated from a certain fungus, was known to suppress T-cell activity, the NEI scientists thought of the possibility that cyclosporine may alleviate this inflammatory eye process. Remarkably the cyclosporine treatment completely prevented the S-antigen induced uveitis in animals.

The laboratory studies encouraged Dr. Nussemblatt to test the effectiveness of cyclosporine in patients with various types of uveitis. Preliminary studies suggested that patients

with the inflammatory disorder known as Behcet's disease were particularly responsive to cyclosporine treatment in that the inflammatory process was dramatically alleviated. These findings served as the basis for a full-scale randomized clinical trial of cyclosporine as a method of treatment of Behcet's patients. Dr. Nussenblatt arranged to have this trial conducted as a multi-center study in Japan where there is greater prevalence of these Behcet's cases than in this country. This study was concluded recently and positive results were obtained indicating that cyclosporine is indeed effective against this form of uveitis. Thus, the series of studies which began in the laboratory has led to the development of a treatment of an inflammatory eye disease where no specific treatment was known before.

For his contributions for the development of cyclosporine as a therapeutic agent of uveitis Dr. Robert Nussenblatt will be honored by the Japanese Ophthalmological Society in 1987.

Dr. Piatigorsky and associates of the LMDB have been developing transgenic mice which have been used in an exciting study that potentially has clinical relevance. These mice permit the analysis of DNA sequences which are responsible for the expression, tissue-specificity and developmental program of gene expression. In their investigations they identified numerous putative regulatory regions of the genes for a specific lens protein, the a A-crystallin. Furthermore, transient expression experiments using explanted lens epithelia have provided evidence that these DNA sequences are indeed responsible for the tissue-specific expression of the crystallin genes. Recently, they created a DNA containing the a A-crystallin gene promoter (only 364 nucleotides) fused with the bacterial chloramphenicol acetyltransferase gene They injected this recombinant DNA into the nucleus of a fertilized mouse egg and produced a male transgenic mouse which had CAT activity in its ocular lens and only in the lens. None of the 9 other tissues examined contained CAT activity. This clearly permits further analysis of the molecular nature of tissue-specific crystallin gene expression at the level of the whole organism. Moreover, it kindles hope for eventual gene therapy of ocular diseases.

For his many important contributions of the application of molecular biology to the eye, Dr. Piatigorsky was the recipient of the 1986 Friedenwald Award, one of the prestigious awards presented by the Association for Research in Vision and Ophthalmology.

Major advances in the study of gyrate atrophy have been made by a team of scientists led by Dr. Muriel Kaiser-Kupfer. Gyrate atrophy is a rare hereditary disease which leads to the degeneration of the retina and choroid. This team of scientists has shown that this disorder is caused by a deficiency of the enzyme ornithine aminotransferase (OAT)

which results in a hyperornithinia in these patients. Dr. Kaiser-Kupfer has shown that diets which restrict the sources of ornithine seem to protect and even improve visual function.

Stimulated by these clinical studies, Dr. George Inana and his colleagues began examining the gyrate atrophy problem using DNA technology. They have been successful in isolating a cDNA clone for human OAT. With this probe they are studying the nature of the OAT gene defect in gyrate atrophy patients. This study is an example of how rapidly a clinical problem can be attacked with the most modern of research tools because of the unique setting of the intramural program which fosters the interactions of scientists from many disciplines.

For the development of the first cDNA probe to study an ocular disease Dr. George Inana was honored as the principal guest lecturer at the Proceedings of the Japanese Chapter of the International Society of Eye Research in Sendai, Japan in 1985.

Dr. Wurtz has pioneered studies on the understanding of how the brain uses visual information to produce eye movements. His recent work, using old world monkeys as a model system, has had two facets. First, he has extended his analyses of the brain circuits that produce the rapid or saccadic eye movements that move the eye quickly from one part of the visual field to another. His studies were the first to recognize that the basal ganglia of the brain participated in controlling saccadic eye movements. The basal ganglia produce a tonic inhibition on the saccadic control system of the brainstem, the superior colliculus. In addition his recent findings revealed that the neuronal transmitter is likely to be GABA since he has been able to inhibit or facilitate eye movements by injecting minute quantities of GABA agonists or antagonists into the terminal area of basal ganglia fibers in the superior colliculus. Thus, a major new control system acting on the initiation of saccadic eye movements has been revealed. His second series of experiments has been on a second type of eye movement, the pursuit eye movements that allow the eye to track moving targets. Dr. Wurtz's group has shown that these movements are normally dependent for their visual input on a tiny area of cerebral cortex that is devoted to processing of visual motion information. Furthermore, minute damage produced by microliter injections of a neurotoxic chemical into adjacent regions produces the directional deficit in pursuit eye movements seen in human patients with cerebral damage. These experiments show for the first time the precise localization and function of the cerebral cortical areas upon which pursuit eye movements are dependent.

For these accomplishments Dr. Wurtz was honored by the European Neuroscience Association which has designated him to present the Gordon Holmes Lecture at the meeting of the Association in September, 1985.

Although the intramural program faces difficult times ahead because of restrictions in budget, personnel and space it is encouraging that accomplishments like those cited here can be achieved. This is a tribute to the intramural scientists with their talents and enthusiasm who devote their lives to dispel our ignorance in many problems related to the eye.

PROJECT NUMBER

Z01 EY 00065-09 OSD

October	f, 1985 to Sep	tember 30, 1	986		
	ECT (80 characters or Wes gical Studies				
PRINCIPAL INVE	STIGATOR (List other pr	ofessional personnel b	elow the Principal Inves	tigator) (Name, title laboratory a	nd institute affiliation)
PI:	Francisco de	Monasterio	M.D., Sc.D	Medical Officer	OSD, NEI
Others:	Edna P. McCr	ane	B.S.	Biologist	OSD, NEI
COOPERATING		rd Modical S	shool Massac	Shugara Fire and Fr	7.6
	lassachusetts.	id Hedical 3	chool, rassac	husetts Eye and Ea	r infirmary,
AB/BRANCH Office of	the Scientif:	ic Director			
SECTION					
NSTITUTE AND	LOCATION				
NEI, NIH,	Bethesda, Mar	vland 20892			
TOTAL MAN-YEA		PROFESSIONAL		OTHER	
0.90		0.40		0.50	
	PRIATE BOX(ES)	_			
	an subjects	(b) Humar	tissues 🕹	(c) Neither	
☐ (a1)	Minors				
□ (a2)	Interviews				

SUMMARY OF WORK (Use standard unreduced type. Do not exceed the space provided.)

This project involves the study of the physiological organization of neurons of the visual system of primates. We have examined the functional mapping on the striate cortex of the innervation density mediated by the two major classes of ganglion cells that respectively project to the parvocellular and magnocellular layers of the lateral geniculate nucleus. We have found that the mannification in striate cortex is proportional to the afferent density of one cell type, and that the so-called point-image area of striate cortex follows the reciprocal of the afferent density of the other cell type. These differences between neural maps are likely to have psychophysical consequences. In addition, an electrophysiological survey of the variation of receptive-field size with eccentricity is in progress, and we have begun preliminary recordings in striate and extrastriate cortex to examine chromatic cell properties. Finally, we are completing analyses of previous studies for their publication.



October 1, 1985 to September 30, 1986

PERIOD COVERED

PROJECT NUMBER

ZO1 EY 00135-14 OSD

	PROJECT (80 characters or less emistry of Retina		ween the borders) ithelium in Health amd Diseas	e
PRINCIPAL	INVESTIGATOR (List other pro	ressional personnel below the	Principal Investigator) (Name, title, laboratory, and instr	tute affiliation)
PI:	Helen H. Hess	M.D.	Medical Officer (Research)	OSD, NEI
				•
COOPERAT	TING UNITS (# any)			
Veteri	inary Resources Br	ranch, DRS, NIH		
AB/BRANC Office	ch e of the Scientif:	ic Director, NEI		
ECTION			-	
	and LOCATION hal Eye Institute	, NIH, Bethesda,	Maryland 20892	
OTAL MAN	N-YEARS	PROFESSIONAL 1.0	OTHER 0.4	
☐ (a) H	PROPRIATE BOX(ES) Human subjects a1) Minors a2) Interviews	(b) Human tissue	es 🔼 (c) Neither	
UMMARY	OF WORK (Use standard unred	luced type. Do not exceed the	space provided)	

The effects of nutrition, oxidation, and other environmental factors (light intensity or darkness) on the incidence and progress of posterior subcapsular opacities (PSO) associated with retinal degeneration are being studied in Royal College of Surgeons (RCS) rats, in which rod photoreceptor outer segment debris accumulates secondary to a phagocytic defect in the retinal pigmented epithelium. Evidence has been obtained that oxidative changes in polyunsaturated fatty acids in the debris lead to water-soluble toxic aldehydes that can be detected in the vitreous, and are toxic to lens membranes. Several diets have been found to prevent mature cataracts, and darkrearing has been shown to prevent the PSO detectable microscopically. By exposing pink-eyed dystrophic rats to constant light of 25 footcandles beginning (1) at 20-23 postnatal days or (2) at birth, we have been able for the first time to demonstrate histopathological changes similar to those in some naturally occurring human posterior subcapsular cataracts (PSC), such as those seen in retinitis pigmentosa. Lens epithelial cells migrated to the posterior pole of the lens. Many were bizarre in shape, with abundant pale-staining cytoplasm and small (or large) nuclei ("bladder cells of Wedl"). Changes similar to human anterior subcapsular cataracts were also RCS rats provide a readily manipulated animal model of PSC, exacerbated by some environmental factors and prevented by others. Principles established with the model may have significance for slowing or preventing human PSC.

ANNUAL REPORT NATIONAL EYE INSTITUTE October 1, 1984 - September 30, 1986

REPORT OF THE DEPUTY CLINICAL DIRECTOR Robert B. Nussenblatt, M.D.

The Clinical Branch consists of three Sections, each with its own Section Head: Section on Ophthalmic Genetics and Pediatric Ophthalmology, Muriel I. Kaiser-Kupfer, M.D.; Section on Neuro-ophthalmology, James Carl, M.D.; and the Section on Retinal and Vitreal Diseases, Robert B. Nussenblatt, M.D. (Acting).

The Section on Ophthalmic Genetics and Pediatric Ophthalmology continues its long-term interest in gyrate atrophy. Patients placed on a low arginine, low protein diet with supplemental amino acids continue to be observed. The group has also been actively pursuing better ways to control the deposition of cystine crystal in the cornea of patients with cyctinosis. Patients with this recessively inherited storage disease will accumulate nonprotein cystine within cellular lyposomes. The ocular manifestations of this disorder include photophobia, crystal deposition in the cornea, conjunctiva, iris, and depigmentation of the retina. A masked randomized clinical trial using topical cysteamine has begun, in order to answer the question of whether this approach will prevent further deposition of crystals in the corneas of cystinosis patients. Another important area of research includes the documentation and monitoring of opacities in the human lens, of great import with the potential availability of medications that may be of use in treating cataract formation.

The Section on Retinal Diseases and Vitreous remains heavily involved with two long term clinical trials. Sorbinil, an aldose reductase inhibitor given orally, is being tested in a randomized masked study to see if it will inhibit the development of diabetic retinopathy. Additionally, patients with senile macular degeneration continued to be studied in this randomized masked study in order to test the efficacy of vitamin E and C therapy as well as the prevention of damage from light below 500 nanometers in preventing this degenerative process, the leading cause of newly registered blindness in the white adult population in the United States. The results of these two most important trials will not be available for some time to come. In addition, this group has studied the amount of flourescein leakage into the vitreous of diabetic patients, using the vitreous fluorophotometer. Diabetic patients with no microaneurysms have not demonstrated greater posterior vitreous leakage as compared to controls. However, differences in vitreal leakage can be noted in diabetic patients with minimal background retinopathy when compared to controls.

The Neuro-ophthalmic Section has concentrated its efforts in two major areas. They have devoted their efforts at developing methods for the analysis of oculomotor disorders in human subjects. This has led to a computerized facility which can stimulate an eye muscre weakness in normals using optical means studies on congenital nystagmous which have shown that these patients

with an abnormality of their vestibulo-ocular reflex, and that some may use this poor response to help them improve acuity by head shaking. Clonazepam has been seen to quiet a variety of types of nystagmus.

The Clinical Branch reflects new horizons with basic research observations playing an increasingly greater role in the research being conducted.

PROJECT NUMBER

Z01 EY 00213-01 CB

PERIOD COVERE					
October	1, 1985 to Septe	mber 30, 1986			
		must fit on one line between the ontributions to o			
PRINCIPAL INVES	STIGATOR (List other professio	na' personnel below the Principa	a' Investigator) (Name title laboratory, ar	id institute affiliation	
PI:	Kent E. Higgins	Ph.D.	Expert	CB, NEI	
Others:	Rafael C. Carus	o M.D.	Visiting Scientist	CB, NEI	
•	Edmond Thall	M.D.	Staff Fellow	CB, NEI	
-	Monique S. Roy	M.D.	Visiting Scientist	CB, NEI	
	Francisco de Mo		Medical Officer		
	Robert Nussenbl		Deputy Director	CB, NEI	
COOPERATING U	NITS (if any)				
None					
LAB/BRANCH					
Clinical	Branch				
SECTION					
Office o	of the Clinical D	irector			
INSTITUTE AND L					
NEI, NIH	I, Bethesda, Mary	land 20892			
TOTAL MAN-YEAR	RS PRO	FESSIONAL	OTHER		
1.	97	1.07	0.90		
CHECK APPROPE	RIATE BOX(ES)				
🛛 (a) Huma	an subjects 🔲	(b) Human tissues	☐ (c) Neither		
₹⊠ (a1)	Minors				
	Interviews				
SUMMARY OF W	ORK (Use standard unreduced	type. Do not exceed the space	provided)		

Spatial contrast sensitivity was used to assess losses or changes in overall visual resolution in patients having a variety of toxic, inflammatory, degenerative, or congenital retinal and neuro-ophthalmological disorders of the visual system. A criterion-free forced-choice psychophysical procedure was used, since this method was previously shown to minimize false positive or false negative diagnoses at initial test and to minimize spurious changes in sensitivity with repeated testing. Contrast sensitivity testing, while requiring more patient testing time, continued to be superior to conventional acuity measuremeths for the detection of early losses and for monitoring changes in visual resolution in patients undergoing treatment. Age-referenced normative data make it possible to distinguish contrast sensitivity loss due to ocular disorder from that expected on the basis of normal aging.

A retinal image stabilization system was used in conjunction with the spatial contrast sensitivity test system to carry our preliminary research on normal subjects using artificial scotomata and complex spatial luminance profiles to examine interactions among fixational errors, eye movements, and visual field defects. This system is currently being modified to permit focal electroretinography and high resolution microperimetry in small, localized regions of the retinas in Eye Clinic patients.

PROJECT NUMBER

Z01 EY 00214-01 CB

PERIOD COVERED October 1, 1985 to September 30, 1986					
Acquired	ECT (80 characters or less Title must fit o and congenital color v	n one line between the ision defici	e borders) encies: Mechanisms and	diagnosis	
PRINCIPAL INV	ESTIGATOR (List other professional person	inel below the Principa	I Investigator) (Name, trite, laboratory, and	institute affiliation)	
PI:	Kent E. Higgins	Ph.D.	Expert	CB, NEI	
Others:	Kenneth B. Knoblauch Edmond Thall Francisco de Monasteri Rafael C. Caruso Robert Nussenblatt	Ph.D. M.D. O M.D. M.D. M.D.	Staff Fellow Staff Fellow Medical Officer Visiting Scientist Deputy Director	CB, NEI CB, NEI CB, NEI CB, NEI CB, NEI	
COOPERATING None	UNITS (# any)				
Clinical	Branch				
SECTION Office of	f the Clinical Director				
NEI, NIH, Bethesda, MD 20892					
TOTAL MAN-YE	1.2 PROFESSION	AL 1.1	OTHER 0.1		
(a) Hun (a1)	Minors Interviews	man tissues	(c) Neither		
SUMMARY OF WORK (Use standard unreduced type. Do not exceed the space provided.)					

This project involves the study of cone function in cases of color vision defects, with special emphasis on the acquired color deficiencies. Human subjects have been used for these studies which range from attempts to improve quantification of data from existing standardized tests of color vision to the collection of additional data for the purpose of designing better tests for detecting color defects secondary to ocular disorder.

DEDARTMENT OF HEALTH A	NO HIMAN CERVICE	C BUBLICHE	I TH SERVICE	PROJE	CT NUMBER	
NOTICE OF INTRAMURAL RESEARCH PROJECT					EY 00117-0	06 CB
PERIOD COVERED October 1, 1985 to Sep	tember 30, 19	86		1		
TITLE OF PROJECT (80 characters or less Oculomotor Disorders i	n Human Subje	ct				
PRINCIPAL INVESTIGATOR (List other prof PI: James R. Carl			ugator) (Name title ad, Section		d institute affiliation	on)
ri. James K. Cari	м	Neuro-opht		СВ	, NEI	
Others: Jon N. Currie	F.R.A.C.P.	Visiting S	Scientist	СВ	, NEI	
Victor Matsuo	Ph.D.	Senior Sta	ff Fellow	CB	, NEI	
COOPERATING UNITS (d any) Laboratory of Sensorim Department of Neurolog					M. Goldbe:	rg);
AB/BRANCH Clinical Branch						
SECTION Section on Neuro-Ophth	almology					
NSTITUTE AND LOCATION NEI, NIH, Bethesda, Ma	ryland 20892					
TOTAL MAN-YEARS	PROFESSIONAL		OTHER			
1.2	1.2		0			
CHECK APPROPRIATE BOX(ES) (a) Human subjects (a1) Minors (a2) Interviews	□ (b) Human tis	ssues 🗆	(c) Neither			

SUMMARY OF WORK (Use standard unreduced type. Do not exceed the space provided.)

The major emphasis of this project has been on developing methods for analyzing oculomotor disorders in human subjects. Further development of the computerized system for stimulus presentation and eye movement recording has enabled us to present brief sets of visual tasks and collect data to evaluate each of the ocular motor subsystems. Additional work on the analysis programs has increased the sensitivity of these tests.

A second phase of the project has been to develop a detailed data base of normal human responses to various stimuli. We have recently concentrated on smooth pursuit eye movements, which have not previously been well characterized.

The computerized facility has allowed us to simulate an eye muscle weakness in normal subjects by optical means, and we found that subjects were able to alter the long latency pursuit response, but not the short latency one, to correct for the simulated weakness. These tests have been applied to a few patients with pursuit disorders and both short and long latency responses were abnormal.

Studies on congenital nystagmus were completed after finding that these patients often have an abnormality of the vestibulo-ocular reflex, and that some may use this poor vestibular response to help them improve acuity by head shaking. Other observations included: benefit of clonazepam in quieting a variety of types of nystagmus, continued analysis of horizontal saccadic abnormalities in all types of Gaucher's disease, and development of abducting nystagmus in multiple sclerosis patients as a result of adaptive changes after eye patching.

PROJECT NUMBER

Z01 EY 00160-04 CB

PERIOD COVERED October 1, 1985 t	o Septer	mber 30, 198	36			
TITLE OF PROJECT (80 characte Visual Inattentic					ons	
PRINCIPAL INVESTIGATOR (List PI: James R	other profession. Carl			igator) (Name title labora ad, Section on		1)
			Neuro-op	hthalmology	CB,	NEI
		F.R.A.C.P.	_		CB,	
Victor	Matsuo	Ph.D.	Staff Fel	low	CE,	NEI
COOPERATING UNITS (# any) Laboratory of Sen M.E. Goldberg, E. LAB/BRANCH Clinical Branch			n, NEI (D.I	. Robinson, S.	E. Peterson,	
SECTION						
Section on Neuro-	ophthalr	nology				
NEI, NIH, Bethesda, 20892						
TOTAL MAN-YEARS	PRO	FESSIONAL		OTHER		
.5		.5				
CHECK APPROPRIATE BOXIES) (a) Human subjects (a1) Minors	_	(b) Human tiss	sues 🗆	(c) Neither		
(a2) Interviews						
BUILDING OF WORK ///co stood		Ama Da mat aucasa	the seems provided	- L		

unreduced type. Do not exceed the space provided.)

Attentional mechanisms important in visual behavior were studied in patients with a variety of central nervous system abnormalities.

Shifts of visual attention as measured by reaction times were measured in patients with parietal lobe damage, and compared to patients with frontal lobe damage, Alzheimer's disease, or schizophrenia.

The patients with parietal lobe dysfunction demonstrated particular difficulty in shifting attention away from the ipsilateral visual field, and this finding was a reliable indicator of parietal cortical dysfunction.

Male patients with idiopathic hypogonadotrophic hypogonadism also had abnormal responses: they were slow in responding to targets in their right visual field.

Eye movements were evaluated in these patient groups, with pursuit movements and fixational stability emphasized. In some patients, square wave jerks present during fixation and reading were indications of attentional disorders.

PROJECT NUMBER

Z01 EY 00162-04 CB

PERIOD COVERED				1
October 1, 1985 to Sep	otember 30, 1986			
TITLE OF PROJECT (80 characters or less	Title must fit on one line between th	e borders)		
Vitreous Fluorophotome	etry			
PRINCIPAL INVESTIGATOR (List other pro		a Investigator) (Name title	e laborator, and institute affination)	
PI: Monique Roy	M D Visitin	- Scientist	CB NEI	
ii. nonique noy		5 50100150	<i>55</i> , 1121	
2.				
COOPERATING UNITS (I' an, I				
None				
None				
LAB/BRANCH				
Clinical Branch				
SECTION				
Section on Retinal and	d Vitneal Diseases			
NSTITUTE AND LOCATION	I Vicreal Diseases			
NEI, NIH, Bethesda, Ma		OTHER		
TOTAL MAN-YEARS	PROFESSIONAL	OTHER		
1.50	1.50	0	.00	
CHECK APPROPRIATE BOX(ES)	□ (b) 11	(a) Na.43		
1	(b) Human tissues	(c) Neither		
(a1) Minors				
(a2) Interviews				
SUMMARY OF WORK (Use standard unred	fuced type. Do not exceed the space	provided)		

Vitreous fluorophotometry has been performed in patients with diabetes mellitus without retinopathy, patients with diabetes mellitus with nonproliferative retinopathy, and normal volunteer subjects, age- and sexmatched to the patients. The amount of fluorescein leakage into the vitreous of patients has been compared to that of the normal subjects. Correlations with other features of diabetes, such as the quality of diabetic control, the existence of subclinical neuropathy and nephropathy, and others were sought.

PROJECT NUMBER

Z01 EY 000198-03 CB

	CT (80 characters or less . Retinopathy T	Title must fit on one line between rial	en the borders)	
PRINCIPAL INVES	STIGATOR (List other prote	ssional personne below the P	rincipal Investigator) (Name title laboratory and in	stitute affiliation)
PI:	Monique Roy	M. D.	Visiting Scientist	CB, NEI
Others:	Manuel Datile James Carl	s M. D. M. D.	Staff Ophthalmologist Senior Staff Fellow	CB, NEI CB, NEI
			nd Metabolic Diseases, Natio	
	itis, Diabetes		nd Metabolic Diseases, Nationand Kidney Diseases, NIH (R	
of Arthr LAB/BRANCH Clinical SECTION	itis, Diabetes Branch		and Kidney Diseases, NIH (R	
of Arthr LAB/BRANCH Clinical SECTION Section INSTITUTE AND L	itis, Diabetes Branch on Retinal and	, and Digestive Vitreal Disease	and Kidney Diseases, NIH (R	
of Arthr Clinical SECTION Section INSTITUTE AND L NEI, NIH	Branch on Retinal and ocation , Bethesda, Ma	, and Digestive Vitreal Disease	and Kidney Diseases, NIH (R	

Oral sorbinil, an aldose reductase inhibitor, will be administered in a double-masked randomized trial to diabetics with no or minimal diabetic retinopathy. This will be done to evaluate the effects of sorbinil on the development of diabetic retinopathy and further investigate the safety and toleration of sorbinil. The study will be conducted simultaneously in 11 research centers in the USA.

PROJECT NUMBER

Z01 EY 00187-03-08

1						1 0 0	9 5
PERIOD COVE October	1, 1985 to Septer	mber 30, 19	986				
	OJECT (80 characters or less Ti ects of Corneal Co						
PRINCIPAL IN	VESTIGATOR (List other profes.	sional personnel bei	low the Princip	al Investigator) (Name title, li	aboratory and institute atti	(ation)	
PI:	Manuel B. Datile	es	M.D.	Visiting Scient	ist	CB,	NEI
Others:	Carl Kupfer		M.D.	Director			NEI
	Lessie McCain		R.N.	Clinical Techni	cian	CB,	NEI
-	Muriel I. Kaiser	r-Kupfer	M.D.	Head, Section o Ophthalmic Ge		CB,	
				•	Ophthalmology		
COOPERATING	G UNITS (if any)						
LAB/BRANCH							
Clinical	Branch						
SECTION	-						
Section	on Ophthalmic Ger	netics and	Pediatri	ic Ophthalmology			
INSTITUTE AN							
NEI, NIH	, Bethesda, Maryl	and 20892					
TOTAL MAN Y	EARS PI	ROFESSIONAL		OTHER .			
. 2		.10		1			
CHECK APPRO	PRIATE BOX(ES)						
🔲 (a1	man subjects) Minors) Interviews	(b) Human	tissues	(c) Neither			
	WORK (Use standard unreduce	ed type Do not exc	eed the space	provided)			
Short- a	s well as long-te	rm effects	of cont	act lens wear o	n the cornea an	re be	eing

Short— as well as long-term effects of contact lens wear on the cornea are being investigated. Changes in corneal curvature, changes in corneal epithelial morphology and changes in corneal endothelial cell morphology are being studied by specular microscopy.

These data will help us understand the dynamics involved in the interaction between a contact lens and the cornea, the risk involved to corneal tissues, and how a systemic or local disorder may increase these risks.

PROJECT NUMBER

Z01 EY 00188-03 CB

PERIOD COVER October		stember 30, 1986		
		Title must fit on one line be toring of Opaci	ties in the Human Lens	
PRINCIPAL INVE	ESTIGATOR (List other pro	dessional personnel below th	e Principal Investigator) (Name, title, laboratory, and i	institute affiliation)
PI:	Manuel B. Dat	iles M.D.	Visiting Scientist	CB, NEI
	Carl Kupfer		Director	NEI
	Robert Sperdu	ito M.D.	Head, Epidemiology Branch	BEP. NEI
-	Peter Kador	Ph.D.	Head, Section on	LMOD, NEI
			Molecular Pharmacology	J. 102 , 112 2
	Lessie McCair	R.N.	Clinical Technician	CB, NEI
Image Pr	ocessing and A	nalysis Laborat	ory, DCRT, NIH (Benes Trus,	Ph.D., Cnief)
Clinical	Branch			
Section Section	on Ophthalmic	Genetics and Pe	diatric Ophthalmology	
NEI, NIH	LOCATION , Bethesda, Ma	ryland 20892		
TOTAL MAN YEA	ARS	PROFESSIONAL	OTHER	
. 7		•5	.2	
(a) Hum	Minors	(b) Human tissu	ues (c) Neither	
	Interviews			
SUMMARY OF V	VUHR (Use standard unred	luced type. Do not exceed th	ne space provided)	

We are developing objective and subjective methods to monitor and document opacities in the human lens using different systems. We are presently actively recruiting patients with and without cataracts for reproducibility studies on the objective systems—the Scheimpflug cameras (Zeiss and topcon), Retroillumination camera (Neitz), Specular microscope (Keeler) and laser light-scattering spectroscope (KOWA). We will also test other systems using sound (ultrasonography), and nuclear magnetic resonance (magnetic resonance imaging). We are also studying subjective systems or method, such as the effects of cataracts on

visual perception, contrast sensitivity, and glare, which may be useful as additional parameters in the monitoring of cataract presence, progression, or

regression.

DC: 14111	MENT OF HEALTH A	IND HUMAN SERV	ICES - PUBLIC HEA	LTH SERVICE		
	NOTICE OF INT	RAMURAL RE	SEARCH PROJE	СТ		
					701 FY	00212-01 CP
PERIOD COVERE	D					
October	1. 1985 to Section 1. 1985 to Se	eptember 30.	1986	e l		
MODEL PI PRINCIPAL INVES	rogram Ior Co. STIGATOR (List other pro	<u>laboration</u> Messiona personnel bi	<u>Between Latar</u> elow the Principa Investi	act Sungeons as gator) (Name, title, laborate	nd Uphtha on and institute	lmic Researc a ^m aton,
	Manuel B. Da			•		
Others:	Carl Kupfer		M.D.	Director		NEI
-	Samuel Zigle	er	Ph.D.	Head, Section Cataracts	n on	
	Peter Kador		Ph.D.	Head, Section Molecular H		•
oconsult	ints'arJin H.	Kinoshita	Ph.D.	Scientific D	irector	NEI
				Head, Section Molecular H	n on	LMOD, NEI
AB BRANCH						
Clinical	Branch					
	on Ophthalmic	Genetics a	nd Pediatric	Ophthalmology		
NSTITUTE AND L			900			
NEI, NIF	i, Bethesda, N	aryland 20	092			
NSTITUTE AND L	i, Bethesda, N RS	anyland 20	092	OTHER		

, PACUECY NUMBER

There is presently an extreme dearth of human cataract material because of an abrupt shift of cataract surgical technique from intracapsular (intact lens) to extracapsular (fragmented lens), primarily because of advent of the use of intraocular lens. We are exploring ways by which fragmented lens materials can be maximally used in cataract basic research through close collaboration

between cataract surgeons and basic researchers and modification of techniques

by both groups.

SUMMARY OF WORK (Use standard unreduced type. Do not exceed the space provided.)

PROJECT NUMBER

Z01 EY 00011-12 CB

PERIOD COVERED							
October 1 1985 to	Sentember 30 1	1986					
October 1, 1985 to	or less. Title must fit on one li	ne between the	borders)				
Pigment Dispersion	With and Withou	ıt Glauco	ma				
PRINCIPAL INVESTIGATOR (List of	her professional personnal bel	ow the Principal	Investigator) (Na.	me, title, laborat	ory, and institute effilia	tion)	
PI: Muriel I.	. Kaiser-Kupfer	M.D.	Geneti	ction on cs and Pe lmology	Ophthalmic diatric	CB,	NEI
Others: Carl Kupf	`er	M.D.	Director				NEI
Lessie Mo		R.N.	Clinical	Technici	an	CB,	
Sandeep J	lain	M.D.	Visiting	Fellow			NEI
COOPERATING UNITS (if eny)							
			· · · · · · · · · · · · · · · · · · ·				
LAB/BRANCH							
Clinical Branch							
SECTION							ĺ
Section on Ophthal INSTITUTE AND LOCATION	mic Genetics and	i Pediatr	ic Ophtha	lmology	· · · · · · · · · · · · · · · · · · ·		
NEI, NIH, Bethesda		92					
TOTAL MAN-YEARS	PROFESSIONAL:	1 25	OTHER.		•		ŀ
1.55		1.35		•	2		
CHECK APPROPRIATE BOX(ES) (a) Human subjects (a1) Minors (a2) Interviews	☐ (b) Human t	issues	☐ (c) Nei	ther			
SUMMARY OF WORK (Use standard	d unreduced type. Do not exce	ed the space p	rovided)				

The purpose of this project is to compare patients with and without glaucoma having pigment dispersion syndrome. The data acquired may enable a determination of the risk of patients with pigment dispersion syndrome to developing glaucoma as well as add to understanding of the pathology of the the disease.

							PROJECT NUMBER	8	
DEPAR	ITMENT OF HEALTH AN	D HUMAN SERVI	CES - PUI	BLIC HEAL	TH SER	VICE		•	
	NOTICE OF INTR	AMURAL RES	SEARCH	PROJE	СТ				
							Z01 EY 0006	52-10	CB
PERIOD COVER	RED						1501 51 000	/ <u>C</u> J V	<u> </u>
October	1. 1985 to Sept	ember 30. 1	986						
TITLE OF PROJ	ECT (80 characters or less T	litie must fit on one l	ine between	the borders)				
Irido-Co	orneal-Endotheli	al (ICE) Sy	ndrome						
PRINCIPAL INVE	ESTIGATOR (List other profes	ssional personnel bel	ow the Print	cipal Investig	ator) (Na	me, title, lab	oratory, and institute affi	iliation)	
PI:	Muriel I. Kais	er-Kupfer	M.D.	Hea	d, Sec	tion o	n Ophthalmic	CB,	NEI
				Ge	enetic	es and	Pediatric		
				01	ohthal	Lmology	•		
Others:	Carl Kupfer		M.D		ector				NEI
	Lessie McCain		R.N.			Techni			NEI
0000000	Manuel Datiles		M.D.	Vis	<u>lting</u>	Scient	ist	CB,	NEI
COOPERATING I	UNITS (# #hy)								
LAB/BRANCH									
<u>Clinical</u> SECTION	Branch								
	0				. 1. 4 1 7				
INSTITUTE AND	<u>on Ophthalmic G</u> LOCATION	enetics_and	Pedia	trie u	onthal	MOTOEA			
NET NIU	I Dathards Man	land 2000	12						
TOTAL MAN-YEA	I <u>Bethesda Mar</u>	YTAIIU ZUDS ROFESSIONAL		10	THER.				
. 35			.25				1		
CHECK APPROP	RIATE BOX(ES)				 ,-		<u> </u>		
(a) Hum	an subjects	(b) Human t	issues		c) Nei	ther			
🗌 (a1)	Minors	, ,			•				
☐ (a2)	Interviews								
SUMMARY OF W	ORK (Use standard unreduc	ed type Do not exce	ed the space	ce provided.)	_				

This project was formerly titled "Progressive Essential Iris Atrophy." Patients are being recruited with progressive essential iris atrophy with or without associated corneal disease. Information is being gathered to evaluate the clinical features and course of the disease process and to investigate aqueous humor dynamics in both affected and unaffected eyes.

PROJECT NUMBER

NOTICE OF INTRAMURAL RESEARCH PROJECT Z01 EY 00083-09 CB PERIOD COVERED October 1, 1985 to September 30, 1986 TITLE OF PROJECT (80 characters or less. Title must fit on one line between the borders.) Gyrate Atrophy of the Choroid and Retina PRINCIPAL INVESTIGATOR (List other professional personnel below the Principal Investigator) (Name, title, laboratory, and institute affiliation) Muriel I. Kaiser-Kupfer M.D. Head, Section on Ophthalmic CB, NEI Genetics and Pediatric Ophthalmology R.N. Clinical Technician CB, NEI Others: Lessie McCain M.D. Visiting Scientist CB. NEI Rafael Caruso CB, NEI Ph.D. Kent Higgins Expert COOPERATING UNITS (# any) The Howard Hughes Medical Institute Laboratory and the Department of Pediatrics, Johns Hopkins University, School of Medicine, Baltimore, Maryland (David L. Valle, M.D.) LAB/BRANCH Clinical Branch SECTION <u>Section on Ophthalmic Genetics and Pediatric Ophthalmology</u> INSTITUTE AND LOCATION NEI. NIH. Bethesda. Maryland 20892 TOTAL MAN-YEARS PROFESSIONAL. OTHER. . 7 1.2 CHECK APPROPRIATE BOX(ES) (a) Human subjects (b) Human tissues (c) Neither (a1) Minors

Patients with gyrate atrophy of the choroid and retina are examined systematically to confirm the diagnosis. Skin fibroblats of affected patients and family members are grown in tissue culture and assayed for ornithine aminotransferase activity. The results will be evaluated for correlation with the presence of homo- or heterozygosity for the disease trait. Patients will be given a trial of pyridoxine to see if serum concentration of ornithine can be reduced, and, if so, the patient will be classified as a "responder," and treatment with pyridoxine will be continued. Nonresponder and responder patients will be placed on a low arginine, low protein, diet with supplemental amino acids and observed for an arrest or improvement of their disease. If patients are not considered eligible for the diet or if they appear unable to comply with the dietary regimen they will be followed to record the natural progress of the condition. Patients with other forms of retinal degeneration, such as retinitis pigmentosa, fundus flavimaculatus, juvenile retinoschisis, are also examined and their courses are compared with gyrate atrophy patients.

(a2) Interviews

SUMMARY OF WORK (Use standard unreduced type: Do not exceed the space provided.)

PROJECT NUMBER

Z01 EY 00163-04 CB

				201 21 00103	04 CD
PERIOD COVERED October 1, 1985 to Se	ptember 30, 198	36			
TITLE OF PROJECT (80 characters or les NIH Interinstitute Me	ss Title must lit on one line l dical Genetics	between the borders Program:	The Genetics	Clinic	
PRINCIPAL INVESTIGATOR (List other p	rolessional personnel below	the Principal Investig	gator) (Name, title, la	boratory, and institute affiliati	01)
PI: Muriel I. Ka	iser-Kupfer M	Ge	d, Section of enetics and phthalmology	Pediatric	CB, NEI
Others: Lessie McCai	n R	.N. Clir	nical Techni	cian	CB, NEI
				•	
COOPERATING UNITS (# @ny)					
Clinical Branch					
Section on Ophthalmic	Genetics and P	ediatric Op	ohthalmology	,	
NEI, NIH, Bethesda, M.	aryland 20892				
TOTAL MAN YEARS	PROFESSIONAL .05		OTHER .1		
CHECK APPROPRIATE BOX(ES) (a) Human subjects (a1) Minors (a2) Interviews	(b) Human tiss	sues 🗆	(c) Neither		
SUMMARY OF WORK (Use standard unit	educed type. Do not exceed	the space provided)		

The Interinstitute Medical Genetics Program and the Genetics Clinic, supported by the Clinical Center, offer a multidisciplinary approach to patients with genetic disease (ZO1 CP 05139-04 CEB). Involved in the program are researchers from all Institutes. Patients evaluated in the clinic reporsent a broad spectrum of genetic disease. During the last year, approximately 423 individuals were seen, representing approximately 100 different disease categories. Due to the high frequency of ocular involvement in many of the cases, almost all the patient were evaluated by Clinical Branch staff of were discussed in consultation. The Clinic serves as a source of interesting case material concerning patients with inherited or developmental abnormalities of the visual system.

In addition to the Genetics Clinic, patients are seen for genetic consultation at the Maryland School for the Blind. This experience has resulted the recruitment of patients into Clinical Branch protocols.

PROJECT NUMBER

Z01 EY 00172- 04 CB

PERIOD COVERE October	:D 1, 1985 to Se	eptember 30,	1986				
	CT (80 characters or less facular Degene		ine betv	veen the	e borders)		
PRINCIPAL INVES	STIGATOR (List other pro	ofessiona personnel bei	on the	Principa	Investigator) (Name title laboratory and	d institute a	"dation
PI:	Muriel I. Ka	iser-Kupfer	М.	D.	Head, Section on Ophthalmic Genetics	CB,	NEI
Others:	Carl Kupfer		м.	D.	Director		NEI
	Monique Roy		М.	D.	Visiting Scientist	CB,	NEI
COOPERATING U None							
LAB/BRANCH							
Clinical	Branch						
Section Section	on Ophthalmic	Genetics an	d Pe	diat	ric Ophthalmology		
NEI, NIH	OCATION , Bethesda, M	D 20892					
TOTAL MAN-YEAR	· -	PROFESSIONAL			OTHER		
	0.43	0.4	3		0.00		
(a) Huma	an subjects Minors	(b) Human	tissue	es	(c) Neither		
	Interviews ORK (Use standard unred)	duced type. Do not ave	and the	FORCE	provided)		
JUNINER COF WI	City (025 21aunain nuis)	ouced type Do not exc	eeu me	share !	provided /		•

This study will determine if patients with severe visual loss because of senile macular degeneration in one eye and with good vision in the second eye can be protected from severe visual loss in the good eye by the administration of vitamin E and vitamin C when exposure of the retina to light below 500 nanometers is diminished. The recruited patients will be randomly assigned either to a treated or an untreated control group and examined at four-month intervals. Follow-up will continue for five years, unless an early beneficial or detrimental effect causes the study to be terminated in less than five years.

PROJECT NUMBER

	NOTICE OF INTRAMURAL RESEARCH PROJECT						00123	-06-0	В
PERIOD COVER October	1, 1985 to Se	otember 3	0, 1986						
TITLE OF PROJECTION	ECT (80 characters or less Psychophysics	Title must fil or of the	Visual Syst	he borders) em				-	
PRINCIPAL INVE	STIGATOR (List other pro	lessional person	nel below the Princi	pal Investigator) (Nama, title, labori	itory and in.	Stitute affiliati	on)	
PI:	Muriel I. Kai	ser-Kupf	er M.D.	Opht	Section on halmic Gen Pediatric	etics	mology	СВ,	NEI
Others:	Rafael C. Car Kent E. Higgi Ralph D. Gunk	.ns	M.D. Ph.D. O.D.	Expert	ng Scienti lmic Physi			CB, CB,	NEI
COOPERATING	UNITS (if any)								
Clinical	Branch								
Section Section	on Ophthalmic	Genetics	and Pediat	ric Opht	halmology				
NEI, NIH	LOCATION , Bethesda, Ma	ryland 2	20892						
TOTAL MAN YEA	ARS	PROFESSION	AL.	ОТН	• 3				
(a1)	RIATE BOX(ES) IAN Subjects Minors Interviews	☐ (b) Hur	nan tissues	□ (c)	Neither				
SUMMARY OF W	VORK (Use standard unrec	luced type Do n	of exceed the space	e provided)					

The visual function of patient with ocular diseases or lesions in the visual pathways and of normal subjects is measured with psychophysical techniques. These data are correlated with those obtained with electrophysiological tests of visual function. The results stained contribute to the diagnosis of ocular and neural disorders that affect vision, and are needed to characterize their nature and evolution. They are also valuable in the assessment of the effect of different forms of treatment on the outcome of these diseases.

PROJECT NUMBER

ZO1 EY 00144-05-CE

			201 EY 00142	1-05-	JB
October	1, 1985 to September 30,	1986			
Clinical	CT (80 characters or less. Life must lit on one l Electrophysiology of the	Visual S	rorders) System		
PRINCIPAL INVE	STIGATOR (List other professional personne tre)	ow the Principa	l Investigator) (Name, title, laboratory, and instituta afficiet	en,	
PI:	Muriel I. Kaiser-Kupfer	M.D.	Head, Section on Ophthalmic Genetics and Pediatric Ophthalmology	CB,	NEI
Others:	Rafael Caruso	M.D.	Visiting Scientist	CB	NEI
	Kent E. Higgins	Ph.D.	Expert	-	NEI
	Doris J. Collie	A.A.	Health Technician		NEI
Clinical	Branch				
Section	on Ophthalmic Genetics an	d Pediatr	ric Ophthalmology		
	ວວຊາວ. , Bethesda, Maryland 208				
.65	PROFESSIONAL . 3	5	• 3		
□ (a2)			(c) Neither		
SUMMENT OF H	and the standard numeradord type. Do not exec	ten tile share b	101000 /		i

The visual function of patients with ocular diseases or lesions in the visual pathways and of normal subjects is measured objectively with electrophysiological techniques. These data are correlated with those obtained with psychophysical tests of visual function. The results obtained contribute to the diagnosis of ocular and neural disorders that affect vision, and are needed to characterize their nature and evolution. They are also valuable in the assessment of the effects of different forms of treatment on the outcome of these diseases.

DEPAR	TMENT OF HEALTI	H AND HUMAN SERVI	CES - PUBL	IC HEALTH SEI	RVICE	PROJECT NUMBER		
		NTRAMURAL RES						
						Z01 EY 0021	1-01 (CB
PERIOD COVER	ED							
October	1, 1985 to 5	September 30.	ne between th	ne borders)				
		rolled Random:			l of To	niosi Cuetosm	ino	
PRINCIPAL INVE	STIGATOR (List other	professional personnel bel	ow the Princip	el Investigator) (N	eme, title, labo	ratory, and institute affilia	tion)	
PI:	Muriel I. P	Kaiser-Kupfer	M.D.	Geneti		n Ophthalmic Pediatric	CB,	NEI
Others	Lessie McCa	d n	R.N.	Clinical	Techni	rian	CB	NEI
0 01101 01	Manuel Dati			Visiting				NEI
COOPERATING I	UNITS (# any)							
	enetics Brandiam Gahl, M.I	ch, NICHD, Nat:	ional In	stitutes o	of Healt)	n, Bethesda, 1	iaryla	and
LAB/BRANCH								
Clinical	l Branch							
SECTION								
Section INSTITUTE AND	<u>on Ophthalmi</u> Locatión	c Genetics and	<u>l Pediat</u>	ric Ophtha	lmology			
NEI. NIE	I. Bethesda.	Maryland 2089	92					
TOTAL MAN-YEA	RS	PROFESSIONAL:		OTHER				
.25		.15				.1		
CHECK APPROP	, ,	(b) Human	icauca	☐ (a) Na	ithor			
	an subjects Minors	(b) Human t	122062	☐ (c) Ne	utilei			

Nephropathic cystinosis is an autosomal, recessively inherited storage disease in which nonprotein cystine accumulates within cellular lysosomes due to a defect in lysosomal cystine transport. Ocular manifestations include photophobia crystal deposits in cornea, conjunctiva iris and depigmentation of the retina. Systemic complications include the Fanconi syndrome, and renal failure.

(a2) Interviews

SUMMARY OF WORK (Use standard unreduced type. Do not exceed the space provided.)

Eight years ago cysteamine, a free thiol which depletes cystine from cells, was introduced in the therapy of cystinotic patients. Although patients had improved growth and stabilized renal function, there was no noticeable effect on the accumulation of corneal crystals. Recent studies showed that corneal cells in tissue culture are readily depleted of cystine by the introduction of cysteamine, making feasible the use of topical ophthalmic cysteamine to circumvent the humoral route. After appropriate animal studies to test for complications which revealed none, we have begun a double-masked clinical trial to test the efficacy of topical cysteamine in humans. Five patients have thus far been enrolled, outcome of the study awaits further observation.

					PROJECT NUMBER	-
DEPAR	TMENT OF HEALTH	AND HUMAN SERVI	CES - PUBL	IC HEALTH SERVICE		
	NOTICE OF IN	ITRAMURAL RES	EARCH P	PROJECT		
					Z01 EY 00084	-08 CB
PERIOD COVER	RED				1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	
October	1. 1985 to S	entember 30. 1	986			
TITLE OF PROJ	ECT (80 characters or le	ss. Title must fit on one li	ne between th	e borders)		
				n Glaucoma or Ocul		
PRINCIPAL INVE	ESTIGATOR (List other p	professional personnel bek	ow the Principa	al Investigator) (Name, title, labor	atory, and institute affilia	tion)
PI:	Carl Kupfer		M.D.	Director		NEI
Others:	Muriel I. Ka	aiser-Kupfer	M.D.	Head, Section on		CB, NEI
•				Ophthalmic Gene		
-				and Pediatric O		
	Lessie McCa:		R.N.	Clinical Technic		CB, NEI
	Manuel B. Da	atiles	M.D.	Visiting Scienti	st	CB, NEI
	Paul Edwards	3	M.D.	Visiting Fellow		CB. NEI
LAB/BRANCH						
<u>Clinical</u>	Branch					
SECTION						
Section INSTITUTE AND	on Ophthalmic	n Genetics and	Pediati	ric Ophthalmology		
		4amland 2000	2			
TOTAL MAN-YEA	l. Belnesda, r ARS	<u>laryland 2089</u> PROFESSIONAL		OTHER.		
0.65		0.55		.1		
CHECK APPROP	BIATE BOX(ES)	1				
🛛 (a) Hum		(b) Human t	issues	(c) Neither		
(a1)	•	<u> </u>		<u> </u>		
, ,	Interviews					
		educed type. Do not exce	and the annua	annual of 1		
SUMMARY OF W	IOHK (Use standard unr	educed type. Do not exce	ed the space	provided.)		
						•
				ting the role of t		
contribu	iting to all o	connective tig	sues ant	erior to the lens	enithelium	the

With recent embryological research indicating the role of the neural crest in contributing to all connective tissues anterior to the lens epithelium, the group of developmental anomalies of the anterior chamber with glaucoma or ocular hypertension is being reviewed.

ANNUAL REPORT NATIONAL EYE INSTITUTE October 1, 1984 - September 30, 1986

REPORT OF THE CHIEF, LABORATORY OF IMMUNOLOGY Robert B. Nussenblatt, M.D.

This past year was the first full year for the Laboratory of Immunology of the National Eye Institute. The year saw the establishment of four Sections within the Laboratory, each with its own Section Head: Section on Clinical Immunology, Alan G. Palestine, M.D., Section on Immunology and Virology, John J. Hooks, Ph.D., Section on Experimental Immunology, Igal Gery, Ph.D., and Section on Immunoregulation, Robert B. Nussenblatt, M.D.

The Section on Clinical Immunology has been particularly interested in new animal models for human intra-ocular inflammatory disease, as well as the role of the neuro-endocrine axis on the immune response. The Section has added insight into the underlying mechanisms of the intra-ocular inflammatory disease produced by endotoxin immunization. Class II antigen expression by nonimmune cells within the eye was noted in the iris and the ciliary body, though no T-cells were present in the inflammatory infiltrate; gammainterferon, a T-cell product, is thought the most potent inducer of class II expression. Though corticosteroid therapy abrogated both the cellular infiltrate and class II expression, only the cellular infiltrate could be prevented when indomethacin was utilized. The role of class II antigen expression on nonimmune cells may play an important role in the localization of the immune response. In separate studies examining the role of pituitary hormones and their effect on the ocular immune response, the Section has noted that modulation of this system appears to have profound effects on the immune response. The concurrent use of bromocriptine, which blocks prolactin production by the pituitary, with a dosage of cyclosporine that ordinarily is not especially effective in preventing the S-antigen induced experimental uveitis model, proved extremely effective, with complete protection affected. These findings will have immediate clinical application, since manipulation of the neuroendocrine system will hopefully permit the use of lower dosages of cyclosporine, and thereby prevent cyclosporine induced nephrotoxicity. These findings suggest that we are just beginning to learn about the possible immunomodulative techniques which should lead to better therapeutic strategies.

The Section on Experimental Immunology has been actively involved in the evaluation of the immune response and how it relates to ocular antigens. The Section has had the opportunity to compare two models of human uveitis induced by two distinct retinally derived antigens, the retinal S-antigen and IRBP. It has been noted that the susceptibility to these diseases is different in lower mammals. Further, delineation of the new IRBP induced model revealed that it was T-cell mediated and that the disease could be actively transferred by lymphocytes. Further the disorder could be induced in monkeys. The disease was a granulomatous uveitis, bearing characteristics similar to such severe human conditions as sympathetic ophthalmia and Vogt-Koyanagi-Harada's disease.

The Section on Immunology and Virology has had a long term interest in the production of T-cell immunomodulators, particularly interferon, and the role these play in ocular structures. The group has recently demonstrated for the first time the presence of IFN-gamma and IL-2 at the site of a localized ocular autoimmune disease. Additionally, the evaluation of exocrine glands involved in Sjogren's syndrome is infiltrated predominantly with T-cells, the majority of which express class II antigens. Moreover, the glandular epithelial cells (ducts and acini) were induced to express these antigens as well. In order to better evaluate the retinal pigment epithelial cell, a cell with immune-like characteristics, the Section has produced the first monoclonal antibody which is directed solely at the human RPE, with studies using this new immune probe just beginning.

The Section on Immunoregulation has been evaluating, in an in vitro system, the role of ocular cells in enhancing an immune response. development of T-cell lines specific to the retinal S-Aq, which are capable of inducing uveitis when transferred to a naive host, has permitted the Section to ask several questions concerning the interaction between immune cells and the eye. A stable long-term rat Muller cell line has permitted the Section to evaluate the interactions between these two pure cell populations. A profound inhibitory effect by these cells on the T-cell has been seen, suggesting that this may be a normal protection mechanism. However, alterations of the Muller cells' membrane can obviate this inhibitory effect. Understanding basic mechanisms in parasitic conditions has also been a major concern of the Section, particularly toxoplasmosis and onchocerciasis. Indeed, in the first immunohistologic study of ocular onchocerciasis, the group has seen that T-cells are the predominant immune cell present and that vascular endothelia, pericytes and fibroblasts in the eye express class II antigens. Studies in the use of cyclosporine for human intra-ocular inflammatory disease has yielded important information concerning the nephrotoxic effects of the drug. Renal biopsies of patients on cyclosporine have demonstrated alterations in renal morphology, supporting the concept that there is a reversible as well as an irreversible component to this secondary effect of the drug. Newer therapeutic strategies, such as the use of bromocriptine combined with cyclosporine as well as new attempts at evaluating the local efficacy of this drug are underway.

The Laboratory of Immunology's first year has produced significant observations both from a clinical and basic research point of view. A better understanding of the basic mechanisms of ocular inflammatory disease is imperative, and the groundwork has now been laid. Further, as with our initial observations with cyclosporine, we sincerely feel that the Laboratory's work will result in better directed and more effective therapies for a group of diseases that are most challenging.



PROJECT NUMBER

Z01 EY 00229-01 LI

October	D 1, 1985 to Septer	mber 30, 1986		
	CT (80 characters or less Title nt of the Size o		ween the borders) duced in Retinal Vessels Using F	PITC-Dextrans
PRINCIPAL INVES	STIGATOR (List other professio	nal personnel below the	Principa Investigator) (Name title laboratory and institute	e affiliation)
PI:	Susan Lightman	M.D.	Visiting Fellow	LI, NEI
Others:	Alan G. Palesti	ne M.D.	Head, Section on Clinical Immunology	LI, NEI
	Einar Stefansson	n M.D.	Visiting Scientist	CB, NEI
LAB/BRANCH				
	rv of Immunology			
SECTION				
	on Clinical Immu	nology		
INSTITUTE AND L	LOCATION			•
NEI, NIH	l, Bethesda, Mary	land 20892		
TOTAL MAN-YEAR	RS PRO	0.5	OTHER 0	-
CHECK APPROPE	an subjects	(b) Human tissu	es 🗵 (c) Neither	
(a1) (a2)	Minors Interviews			
, , ,	ORK (Use standard unreduced	type Do not exceed the	e space provided)	

Uveitis was induced in two monkeys by immunization with IRBP and serial fluorescein angiograms performed using different sized dextrans linked to fluorescein. The aim of these studies is to provide data on the retinal vessels and toxicology data to enable these agents to be used in humans. We have demonstrated that the larger molecular weight dextrans are less permeable than sodium fluorescein in the inflamed retina.

PROJECT NUMBER

Z01 EY 00230-01 LI

PERIOD COVERED			•					
October 1, 1985 to September 30, 1986								
TITLE OF PROJECT (80								
Quantitativ	e Assessmen	t of Reti	nal Vascu	ılar Pe	ermeability	7		
PRINCIPAL INVESTIGAT	TOR (List other prof.	essiona personne	below the Prin	cipal Investi	igator) (Name_title	laboratory an	id institute affilia	ation;
PI:	Susan Ligh	itman	M.D.	Visiti	ing Fellow			LI, NEI
Others:	Alan G. Pa	lestine	M.D.		Section or unology	n Clinica	al	LI, NEI
Laboratory (M.D.); Labo Rapoport, M	of Neurosci ratory of N	lences, Na Neuroscien	tional Ir ces, Nati	nstitut ional l	te on Aging Institute (g (Emanue on Aging	el Rechtl (Stanle	hand, y
Laboratory	of Immunolo	gy						
SECTION Section on	Clinical In	nmunology						
NEI, NIH, B	-	aryland 2	0892					
TOTAL MAN-YEARS 0.08		PROFESSIONAL 0.	08		OTHER	0		
CHECK APPROPRIATE (a) Human su (a1) Mino (a2) Inten	ibjects rs	(b) Hum	an tissues	Æ	(c) Neither			
SUMMARY OF WORK (UMMARY OF WORK (Use standard unreduced type. Do not exceed the space provided.)							

A sensitive quantitative method was set up for examining the permeability of retinal vessels in the rat. Baseline values for normal rat retinal vessels were established and the method will be applied to pathological situations.

PROJECT NUMBER

Z0I EY 00217-01 L1

PERIOD COVE	RED 1, 1985 to September 30, 1986			
	JECT (80 characters or less Title must fit on one line te Migration in Experimental			
PRINCIPAL INV	ESTIGATOR (List other professional personnal below	the Principal Invest	igator) (Name title, laboratory, and institute i	affiliation)
PI:	Alan G. Palestine	M.D.	Head, Section on Clinica	al LI, NEI
			Immunology	
Others:	Robert B. Nussenblatt	M.D.	Deputy Clinical Director	NEI
:	Consuelo Muellenberg-Coulomb	re	Chemist	LI, NEI
	Myung Kim	M.D.	Visiting Fellow	LI, NEI
	Susan Lightman	M.D.	Visiting Fellow	LI, NEI
			_	
COOPERATING	UNITS (d any)			
Laborato	ry of Immunology			
SECTION	Ty of Imagenotogy			
	on Clinical Immunology			
INSTITUTE AND				
NEI, NIH	, Bethesda, Maryland 20892			
TOTAL MAN-YE			OTHER	
0	.3 0.3		0	
	PRIATE BOX(ES)			
	man subjects \square (b) Human tiss	sues 🖾	(c) Neither	
	Minors			
	Interviews			
CLIMANADY OF	MODE III a secondard consequence and a secondary	*** *** *** ***		

SUMMARY OF WORK (Use standard unreduced type. Do not exceed the space provided.)

Experimental autoimmune uveitis (EAU) is induced by immunization of rats and other experimental animals with S-antigens (a soluble antigen from the retina) is being investigated in this laboratory as a model of human intra-ocular inflammation. This experimental inflammation can be transferred from donor rats to naive recipients using lymphocytes harvested from the spleen or lymph nodes. Following harvesting of the cells from the donors and three days in culture with stimulating antigen, the cells are injected into the intra-peritoneal cavity and five to seven days later the recipient rats develop EAU. The disease can also be transferred using a T-helper cell line by intra-peritoneal or intra-ocular injection. The mechanism of transfer of disease is unclear. This work has used radioactively labeled lymphocytes to determine the fate of these lymphocytes after injection into the peritoneal cavity or blood during the process of the development of uveitis. The goal of this project is to understand the initiating mechanisms of inflammation in the hope that these mechanisms can be extended and applied to human inflammations. Thus far we have determined that only a small percentage of the lymph node lymphocytes injected into the peritoneal cavity actually reach the eye during the induction of EAU. Of 100 million cells transferred into the peritoneal cavity, approximately 5,000 reach the eye. Many more reach the spleen, liver and thymus however. The work is now being extended to study the migration patterns of intra-peritoneal or intra-ocular T-cell lines.

PROJECT NUMBER

Z01 EY 00218-01 LI

PERIOD COVER	ED					
October 1	, 1985 to S	eptember 30, 1	986			
		r less. Title must fit on one		ne border	s)	
Human T-c	ell Leukemi	a/Lymphotropic	Virus Ty	ype I	II and Eye Disease	
PRINCIPAL INVE	STIGATOR (List offer	er professional personnel be	Now the Principa	al Investi	gator) (Name, title, laboratory, and institu	ite affiliation)
PI:	Alan G. Pa	lestine	M.D.		, Section on Clinical munology	LI, NEI
Others;	Leslie S. Robert B. Myung Kim	Fujikawa Nussenblatt	M.D. M.D. M.D.	Depu	or Staff Fellow ty Clinical Director ting Fellow	LI, NEI NEI LI, NEI
		_			ogy, National Cancer I	
			•		& Molecular Biology,	
				-	rtment of Critical Car of Tumor Cell Biology	
LAB/BRANCH			Cancer	r Ins	titute (Robert C. Gall	o, M.D.)
Laborator	y of Immuno	logy				
SECTION						
Section o	n Clinical	Immunology				
INSTITUTE AND						
NEI, NIH,	Bethesda,	Maryland 2089	2			
TOTAL MAN-YEA		PROFESSIONAL			OTHER	
0.9	91	0.	91		0	
	an subjects Minors Interviews	□ (b) Human			(c) Neither	
SUMMARY OF W	ORK (Use standard	unreduced type. Do not ex	ceed the space	provided)	
possible		nt in ophthalmi			s in order to characte AIDS and possible tra	
Schi	rmer's filt		s and inf		of patients with AIDS on of lymphocytes plus	-
					al cells from AIDS pa s a reservoir for the	
		suggest more would be suggest work with a suggest more with a suggest to be suggested as a suggest to be suggested as a suggest to be suggested as a suggest more with			esence of HTLV-III in ous.	

Patients with AIDS and cytomegalovirus retinitis were studied to improve therapy for this blinding disorder. Laser therapy for active lesions was ineffective,

but the antiviral drug DHPG was effective in treating but not curing the

infection.

PERIOD COVERED

PROJECT NUMBER

Z01 EY 00219-01 LI

October	1, 1985 to Sep	tember 30, 1986				
TITLE OF PROJ	ECT (80 characters or less	Title must fit on one line between	the border	s)		
		iptine on Human Uve				
PRINCIPAL INVE	STIGATOR (List other pro	dessional personnel below the Prin	cipal Invest	igator.) (Name, title, laboratory, and institute affiliation	(חי	
PI:	Alan G. Pales	stine	M.D.	Head, Section on Clinical Immunology	LI,	NEI
Others:	Consuelo Muel	llenberg-Coulombre		Chemist	LI,	NEI
-	Myung Kim		M.D.	Visiting Fellow	LI,	NEI
	Robert B. Nus	ssenblatt	M.D.	Deputy Clinical Director		NEI
COOPERATING	UNITS (# any)					
Metaboli	sm Branch, Nat	ional Cancer Insti	tute (Marie C. Gelato, M.D.)		
1						
LAB/BRANCH	6.7					
	ory of Immunolo	ogy				
SECTION		1				
Section INSTITUTE AND	on Clinical In	munology				
		20002				
TOTAL MAN-YE	I, Bethesda, Ma	PROFESSIONAL:		OTHER		
	. 29	0.29		0		
CHECK APPRO	PRIATE BOX(ES)					
		☐ (b) Human tissues		(c) Neither		
· · · ·	Minors	_ (5)	_			
	Interviews					
		duced type. Do not exceed the spa	ce provide	d.)	•	

In recent years there has been increasing evidence in the literature that pituitary hormones are capable of regulating the immune system. There is evidence to suggest that prolactin is an immunostimulatory hormone and that reduction of serum prolactin levels in experimental animals by hypothesectomy or bromocriptine will result in a degree of immunosuppression.

This information has been applied to humans and two clinical studies have begun. Both of these are in early phase of patient recruitment. One study is a randomized trial between placebo and bromocriptine in recurrent anterior uveitis using the end point of the number of recurrences per year to determine whether or not bromocriptine is capable of regulating the immune system in these patients. The second trial focuses on the additive effects of cyclosporine plus bromocriptine in attempts to treat patients with posterior uveitis at lower doses of cyclosporine in order to reduce its concurrent renal toxicity while at the same time achieving an immunosuppressive effect. Cyclosporine and prolactin compete for binding sites on the lymphocyte.

Further studies in human disease will hopefully elucidate other aspects of the neuroendocrine axis which can be utilized to regulate the immune system to treat autoimmune diseases.

DEPARTMENT OF HEALTH AND HUMAN SERVICES - PUBLIC HEALTH SERVICE

PROJECT NUMBER

201 FY 00220-01 IT

	NOTICE OF INTRAMURAL RESEARCH	PROJEC	T 301 21 0022	5 01 L1
PERIOD COVERE October	0 1, 1985 to September 30, 1986			
TITLE OF PROJECTION	CT (80 characters or less Title must lit on one line between the Modulation of Immune-Mediated E	ye Dis	ease in Rats	
PRINCIPAL INVES	STIGATOR (List other professional personnel below the Princip	oal Investige	ator) (Name, title, laboratory and institute affiliat	tion)
PI:	Alan G. Palestine	M.D.	Head, Section on Clinica Immunology	l LI, NEI
Others:	Consuelo Muellenberg-Coulombre		Chemist	LI, NEI
-	Myung Kim	M.D.	Visiting Fellow	LI, NEI
	Robert B. Nussemblatt	M.D.	Deputy Clinical Director	NEI
LAB/BRANCH				
	ry of Immunology			
SECTION	· · · · · · · · · · · · · · · · · · ·			
Section	on Clinical Immunology			
INSTITUTE AND L				
	, Bethesda, Maryland 20892			
TOTAL MAN-YEAR 0.1		0	0 0	
CHECK APPROPR	<u> </u>	·	.	
(a) Huma	· · · · · · · · · · · · · · · · · · ·	₩ (c) Neither	
☐ (a1) M	nterviews			
. , ,	ORK (the standard unreduced hore. Do not exceed the space			

In recent years there has been increasing evidence in the literature that pituitary hormones are capable of regulating the immune system. There is evidence to suggest that prolactin is an immunostimulatory hormone and that reduction of serum prolactin levels in experimental animals by hypothesectomy or bromocriptine will result in a degree of immunosuppression.

An animal model of experimental autoimmune uveitis (EAU) induced by immunization of rats with S-antigen (a soluble antigen from the retina) is used as a model for intraocular inflammatory disease. We have demonstrated that concurrent antibody production in both males and females and the incidence of uveitis in female animals but did not have a significant effect on the immune responses measured by lymphocyte proliferation. As reported before, cyclosporine in high doses (10 mg/kg) there is only partial effect. We have demonstrated that the concurrent use of bromocriptine to suppress prolactin in combination with low dose cyclosporine is more effective than either drug separately in suppressing both the incidence of disease as well as the cellular and humoral immune responses to immunization. There is evidence in the literature to suggest that cyclosporine is able to compete for binding on the lymphocyte by prolactin and that reductions in prolactin level may therefore make cyclosporine more effective. Further studies in animal disease will hopefully elucidate other aspects of the neuroendocrine axis which can be utilized to regulate the immune system to treat autoimmune diseases.

October 1, 1985 to September 30, 1986

PERIOD COVERED

PROJECT NUMBER

Z01 EY 00221-01 LI

		s. Title must fit on one line between th				
Intraocula	r Class II A	ntigen Expression in	Endot	toxin-Induced Uveitis		
PRINCIPAL INVEST	IGATOR (List other pro	ofessional personnel below the Princip	_	ator) (Name, title, laboratory, and institute affiliation		
PI:	Alan G. Pal	estine	M.D.	Head, Section on Clinical Immunology	LI,	NEI
Others:	Myung Kim		M.D.	Visiting Fellow	LI,	NEI
-	Consuelo Mu	ellenberg-Coulombre		Chemist	LI,	NEI
	Robert B. N	_	M.D.	Deputy Clinical Director		NEI
COOPERATING UN	ITS (if any)					
LAB/BRANCH						
	of Immunolog	<u>gy</u>				
SECTION	01					
	Clinical Imm	munology				
INSTITUTE AND LO		1 1 20000				
		ryland 20892		77.150		
TOTAL MAN-YEARS		PROFESSIONAL	1	OTHER		
0.5		0.51		0		
CHECK APPROPRIA				7 A AL 201		
(a) Humar		(b) Human tissues	X	(c) Neither		
☐ (a1) M						
☐ (a2) In						
		duced type. Do not exceed the space				
				cell wall of gram negative		
bacteria.	When inject	ted into the footpad	or th	he eve of a rat it will ind	iuce	an

inflammatory reaction within the eye. Endotoxin injection in the footpad does not lead to inflammation in other organs but does produce bilateral uveitis primarily centered at the ciliary body and iris. The mechanism of this inflammation is still unclear. However, since several types of anterior uveitis in humans appear to be linked to gram negative bacteria exposure, this is considered a relative model for anterior uveitis in humans such as Reiter's syndrome. In this study rats received E. coli endotoxin and the expression of class II antigens was studied within the eye using immunohistochemical techniques. We observed that the expression of class II antigens on the ciliary body and iris preceded the influx of inflammtory cells into the eye and that the inflammatory cells that entered the eye were primarily neutrophils with some monocytes. No T-cells were present in the inflammatory infiltrate. The expression of class II antigens within the eye was restricted to the iris and ciliary body during this inflammatory episode and did not involve the retinal pigment epithelium or retinal vessels. The inflammatory cellular infiltrate could be inhibited by indomethacin or colchicine, however this did not alter the expression of class II antigens by the iris or ciliary body indicating that this expression is not simply a consequence of the inflammatory infiltrate but may be intimately involved with the mechanism of the expression of endotoxin induced uveitis. Corticosteroids were capable of suppressing both the cellular inflammatory infiltrate and the expression of class II antigens. The expression of class II antigens on nonlymphoid cells within the eye may be important in antigen presentation or may simply signal a phenotypic change on the cells due to the interaction of endotoxin with the cell membranes.

PROJECT NUMBER

Z01 EY 00231-01 LI

PERIOD COVERE	D							_
October 1	, 1985 to Sept	ember 30	, 1986					
TITLE OF PROJE	CT (80 charecters or less	Title must fit on	one line between	the borders	;)			
Cell Surf	ace Antigens o	n Retino	blastoma (Cells				
PRINCIPAL INVES	STIGATOR (List other prof	essional personi	nel below the Prin	cipal Investig	gator)	(Name, title, laboratory and institute affil	ation)	
PI:	Barbara Detri	ick Ph	.D.	Ex	pert	t	LI, NEI	ĺ
Others:	John J. Hooks	Ph	.D.			Section on Immunology Virology	LI, NEI	[
-	Gerald J. Cha	der Ph	.D.	Ch	ief		LVE, NEI	I
	Merlyn Rodrig	gues M.	D., Ph.D.			Section on Clinical Pathology	LOP, NEI	[
	Chi-Chao Chan	M.	D.	St	aff	Ophthalmologist	LI, NEI	Ī
COOPERATING U	INITS (if any)							
Duke Univ	ersity, Durham	, North	Carolina (Barton	F.	Haynes, M.D.)		
LAB/BRANCH								-
Laborator	v of Immunolog	v						
SECTION								
Section o	n Experimental	Immunol	ogv					
INSTITUTE AND L	OCATION			-				
NEI, NIH,	Bethesda, Mar	yland 2	0892					
TOTAL MAN-YEAR	RS	PROFESSIONA	AL		OTHE	R		
	0.75		0.75			0		
CHECK APPROPE				_				
🔽 (a) Huma	an subjects	🗵 (b) Hun	nan tissues		(c) I	Neither		
🔲 🔲 (a1) l	Minors							
☐ (a2) I	Interviews							
SUMMARY OF WO	DRK (Use standard unredu	uced type. Do n	of exceed the spa	ce provided)			

Class II antigens (HLA-DR and HLA-DQ) are membrane bound glycoproteins encoded by genes in the major histocompatibility complex. In addition to their well established role as regulatory molecules of the immune response, these determinants are now suspected of playing an influencial part in cellular differentiation.

In exploring the cellular composition of a popular childhood tumor, retinoblastoma, we have had an opportunity to describe class II antigens on a population of undifferentiated malignant cells of the retina. This study provides the initial description of class II antigens on retinoblastoma cells. Furthermore, HLA-DR antigen was found to be coexpressed on cells that contained both neuronal and glial markers. This study also identifies for the first time the présence of class II antigens on cells of neuronal origin.

Based on these initial studies, two areas will be explored. The first approach will focus on the possible role of class II antigens in the cellular differentiation or immune reactivity. The second will examine the prognostic significance of these molecules on retinoblastoma cells and the possible relationship class II proteins may have to the modulation and management of this tumor.

PROJECT NUMBER

Z01 EY 00235-01 LI

PERIOD COVER				
October	1, 1985 to Septer	mber 30, 19	86	
	ECT (80 characters or less. Title			
Identifi	ication and Modula	ation of Cl	ass II Antigens	
			the Principal Investigator) (Neme, title laboratory, and instr	
PI:	Barbara Detrick	Ph.D.	Expert	LI, NEI
Others:	John J. Hooks	Ph.D.	Head, Section on Immunology	LI, NEI
			and Virology	
	Chi-Chao Chan	M.D.	Staff Ophthalmologist	LI, NEI
	Leslie Fujikawa	M.D.	Senior Staff Fellow	LI, NEI
	Richard Wetzig	M.D.	Senior Staff Fellow	LI, NEI
	Caroline Percopo	A.B.	Biologist	LI, NEI
	Paris, France (Lau		, Durham, North Carolina (Barton sell, M.D.): and Paris, France (•
Laborato	ory of Immunology			
SECTION				
Section	on Experimental	Immunology		
INSTITUTE AND	LOCATION			
NEI, NIE	l, Bethesda, Marvi	land 20892		
TOTAL MAN-YEA	ARS PRO	FESSIONAL	OTHER	-
	0.6	0.5	0.1	
CHECK APPROP	_			
(a) Hum	•	(b) Human tis	ssues \square (c) Neither	
_ ` '	Minors			
	Interviews			
SLIMMARY OF W	VORK Illse standard unreduced	Nos Do not exces	The space provided)	

Class II antigens are membrane bound glycoproteins that are encoded by genes in the mixed histocompatibility complex (MHC). The expression of these antigens allows a cell to participate in the initiation and perpetuation of immune responses. Furthermore, although most cells that constitutively express class II antigens are members of the immune system, other cell types can be induced to demonstrate these molecules under selected conditions, such as an immunologic or degenerative event. Early investigations demonstrated that patients with the retinal degenerative disorder retinitis pigmentosa had a unique alteration in two regulatory proteins, interferon-gamma and the MHC class II antigen, HLA-DR. In addition, in vitro studies revealed that a regulatory cell in the eye, the rpe cell, could also express this antigen and that these determinants were sensitive to modulation with interferon-gamma. Based on these findings we expanded our studies to evaluate class II antigen expression in ocular diseases. We found that the rpe cell does not express class II antigen in the normal eye. In contrast, the rpe cell did express these molecules in a retinal degenerative disorder (retinitis pigmentosa) and in two ocular inflammatory diseases (sympathetic ophthalmia and uveitis). Using the EAU animal model of ocular autoimmune disease we demonstrated that the rpe cell is activated to express class II antigens prior to clinical and histopathological evidence of the disease. We are now evaluating the effects of such modulators as interferon-gamma, anti-Ia antiserum and cyclosporine on class II antigen expression with the hope that an alteration in activation or expression of these molecules may modify the disease process to the benefit of the host. In summary, these studies indicate that the appearance and modulation of class II antigens may play a role in the initiation, maintenance or regulation of pathologic events in degenerative or inflammatory processes.

PROJECT NUMBER

Z01 EY 00069-09 LI

october 1, 1985 to September 30, 1986									
INTLE OF PROJ Immune Re	ECT (80 characters or less sponses to ocu	Title must fit o	on one line betwee gens	n the borde	rs)				
PRINCIPAL INV	ESTIGATOR (List other pro	tessional perso	nnel below the Pni	ncipal Inves	tigator) (Nami	e title la	boratory and institute	affiliation)	
PI:	Igal Gery		Ph.D.	-	Section nology	on E	xperimental	LI,	NEI
Others:	Shigeto Hiro Cathy McAlli Barbara Vist Gregory Fox	ster 1	Ph.D. B.A.	Extram Microb	ng Fello ural Fe iologis IH Rese	llow t	Scholar	LI, LI, LI,	NEI NEI
COOPERATING Tokyo Med	UNITS (rany) lical College H	ospital,	Tokyo, Ja	pan (D	r. M. U	sui)			
Laborator	y of Immunolog	у							
Section o	n Experimental	Immunol	ogy						
NEI, NIH,	Bethesda, Mar	yland 20	892						
TOTAL MAN-YE	ARS 2.7	PROFESSION	2.1		OTHER		0.6		
(a) Hun (a1) (a2)	PRIATE BOX(ES) nan subjects Minors Interviews		man ti ssu es		(c) Neit	her			
SUMMARY OF This	WORK (Use standard unred project is all	duced type Do ned at le	not exceed the spearning ab	ace provide Out th	_{d)} e patho	genes	is of immune	-media	ated

eye diseases, mainly by investigating the animal disease, "experimental autoimmune uveitis" (EAU), which is induced in various animals by immunization with certain eye components. EAU is considered a model for certain human ocular conditions. Research during FY 1986 has focused on EAU induced by interphotoreceptor retinoid-binding protein (IRBP). Our previous study (FY-1985) showed that IRBP is highly uveitogenic in Lewis rats. Major new findings include: (1) Rats of different inbred strains vary in their susceptibility to EAU induced by IRBP or by another retinal protein, S-antigen (S-Ag). This finding shows that the susceptibility to EAU is related to genetic makeup and supports the assumption that the susceptibility to immune-mediated diseases in man is also genetically regulated. (2) The pathogenic mechanism of IRBP-induced EAU was found to be cell mediated: the disease could be adoptively transferred by lymphocytes and a correlation was found between EAU development and cellular immunity but not with antibody production. (3) Monkey IRBP was found to be 20 times less uveitogenic in rats than bovine IRBP. The two IRBPs showed cross reactivity when used for stimulation of lymphocytes for EAU induction, but did not cross react by the lymphocyte proliferation assay. This finding indicates that lymphocyte responses in vitro do not necessarily represent their capacity to induce disease in vivo. (4) Monkeys were found highly susceptible to IRBPinduced EAU. This monkey disease is of special interest by showing a close similarity to certain ocular diseases in man, in particular sympathetic ophthalmia and Vogt-Koyanagi-Harada disease. Similarly to these human diseases, IRBP-induced in monkey was expressed mainly as granulomatous choroiditis. In addition to providing a useful model for the human diseases, the findings with IRBP-induced EAU in monkeys support the notion that autoimmune processes to retinal antigens participate in the etiology of certain human eye diseases.

PROJECT NUMBER

Z01 EY 0023-08 LI

PERIOD COVE								
October 1, 1985 to September 30, 1986								
	DJECT (80 characters or less			,				
Macropha	ge and Lymphocyt	e Participa	tion in Infl	ammatory Processes				
PRINCIPAL IN	VESTIGATOR (List other prof	essional personnel be	alow the Principal Inves	igator) (Name title laboratory and	institute effiliation,			
PI:	Igal Gery	Ph.D.	Head, Sect Immunolo	ion on Experimental	LI, NEI			
Others:	Cathy McAlliste	r Ph.D.	Extramural	Fellow	LI, NEI			
	Barbara Vistica	B.A.	Microbiolo	gist	LI, NEI			
Laborato	GUNITS (f any) ny of Developmer nd Human Develop			ity, National Insti	tute of Chile	d		
LAB/BRANCH								
Laborato	ry of Immunology	,						
SECTION								
Section	on Experimental	Immunology						
INSTITUTE AN	D LOCATION							
NEI, NIH	, Bethesda, Mary	land 20892						
TOTAL MAN-Y	EARS	PROFESSIONAL		OTHER				
	0.1			0.1				
	OPRIATE BOX(ES)							
· <u></u> -	•	😠 (b) Human	tissues L	(c) Neither				
_ `) Minors							
) Interviews							
SUMMARY OF	WORK Illse standard unred.	red time Do not ev	services are necessaria	√]				

ORK (Use standard unreduced type. Do not exceed the space provided.)

This project has been aimed at collecting information concerning the processes' which bring about inflammation and the involvement of lymphocytes, macrophages and immune mediators in these processes. Experiments carried out this year have further examined the effects of pertussis toxin (PT) on lymphoid cells in culture. This bacterial product is a powerful adjuvant and it facilitates remarkably the induction of experimental autoimmune diseases. Our previous experiments (FY 1985) have shown that PT stimulates lymphocytes and macrophages in culture. Experiments carried out this year have indicated that PT has a dual effect on these lymphoid cells. Thus, cultures of lymphocytes or macrophages which are stimulated by other agents are often inhibited by the addition of PT at doses which are simulatory to cultures with no other stimuli. This finding is in line with reports showing that, at certain circumstances, PT may both enhance and inhibit immune responses.

This project is being phased out, in order to focus the Section's effort on issues which relate directly to the immunopathogenesis of ocular inflammatory diseases (see project # 201 EY 00069-09).

PROJECT NUMBER

Z01 EY 00232-01 LI

PERIOD COVE				
October	1, 1985 to September	30, 1986		
	JECT (80 characters or less. Title mu			
Interfer	on System in Cellula	ar Functio	n and Disease	
PRINCIPAL INV	ESTIGATOR (List other professional	personnel below	the Principal Investigator) (Name, title Taboratory, and institute	affiliation)
PI:	John J. Hooks	Ph.D.	Head, Section on Immunology and Virology	LI, NEI
Others;	Barbara Detrick	Ph.D.	Expert	LI, NEI
	Caroline Percopo	A.B.	Biologist	LI, NEI
	Yotanna Dalavanga	M.D.	Visiting Fellow	LI, NEI
	Chi-Chao Chan	M.D.	Staff Ophthalmologist	LI, NEI
LAB/BRANCH				
	ry of Immunology			
SECTION	on Immunology and Ud	w. l		
INSTITUTE AND	on Immunology and Vi	rology		
	, Bethesda, Maryland	20892		
TOTAL MAN-YE	PROFE 1.45	SSIONAL 0.75	OTHER 0.7	
(a) Hur	PRIATE BOX(ES) man subjects	Human tiss	sues 🗵 (c) Neither	

SUMMARY OF WORK (Use standard unreduced type. Do not exceed the space provided.)

The IFN proteins can modify a variety of biological activities and are considered one of the body's regulatory proteins. Numerous studies now indicate that the IFN's are potent immunoregulators. During the past year we have been studying the ways in which IFN proteins interact with cells of the immune system and how this interaction may modify immune responses and immunologically related disorders.

Using immunocytochemical analysis we have developed a sensitive method of identifying lymphokines, IFN-gamma and IL2, at the site of tissue damage. We have identified the lymphokines, IFN-gamma and IL2 in inflammatory eye diseases. The presence of these lymphokines is associated with a lymphocyte infiltrate predominantly of a T-cell origin and with the expression of MHC class II antigens on both the infiltrating cells and in the retinal pigment epithelial (rpe) cells.

This is the first demonstration of lymphokines, IFN-gamma and Il2 at the site of a localized autoimmune disease. These observations may indicate that IFN-gamma induced MHC class II antigen expression may serve as a local amplification system in autoimmune and inflammatory eye disease. A better understanding of the role of lymphokines in the mechanisms involved in the development of autoimmunity and inflammation may be beneficial in the treatment of these diseases.

PROJECT NUMBER

Z01 EY 00232-01 LI

PERIOD COVER	ED			
October 1	, 1985 to Septe	ember 30, 1986		
TITLE OF PROJE	CT (80 charecters or less	Title must fit on one line be	tween the borders)	
Interfero	n System in Cel	lular Function	and Disease	
PRINCIPAL INVE	STIGATOR (List other profe	ssional personnel below the	Principal Investigator) (Name title laboratory, and ins	titute affiliation)
PI:	John J. Hooks	Ph.D.	Head, Section on Immunology and Virology	LI, NEI
Others;	Barbara Detric	k Ph.D.	Expert	LI, NEI
_	Caroline Perco	po A.B.	Biologist	LI, NEI
	Yotanna Dalava	inga M.D.	Visiting Fellow	LI, NEI
	Chi-Chao Chan	M.D.	Staff Ophthalmologist	LI, NEI
COOPERATING I	UNITS (d any)			
New York	University, Sch	ool of Medicin	e, Department of Microbiology	(Jan Vilcek,
M.D.)				
LAB/BRANCH				
	y of Immunology	,		
SECTION				
Section o	n Immunology ar	nd Virology		
INSTITUTE AND	LOCATION			
NEI, NIH,	Bethesda, Mary	land 20892		
TOTAL MAN-YEA	RS	PROFESSIONAL	OTHER	
1	. 45	0.75	0.7	
CHECK APPROP	RIATE BOX(ES)			
🔲 (a) Hum	an subjects	🗌 (b) Human tissu	ies 🛭 (c) Neither	
☐ (a1)	Minors			
(a2)	Interviews			

SUMMARY OF WORK (Use standard unreduced type. Do not exceed the space provided.)

The IFN proteins can modify a variety of biological activities and are considered one of the body's regulatory proteins. Numerous studies now indicate that the IFN's are potent immunoregulators. During the past year we have been studying the ways in which IFN proteins interact with cells of the immune system and how this interaction may modify immune responses and immunologically related disorders.

Using immunocytochemical analysis we have developed a sensitive method of identifying lymphokines, IFN-gamma and IL2, at the site of tissue damage. We have identified the lymphokines, IFN-gamma and IL2 in inflammatory eye diseases. The presence of these lymphokines is associated with a lymphocyte infiltrate predominantly of a T-cell origin and with the expression of MHC class II antigens on both the infiltrating cells and in the retinal pigment epithelial (rpe) cells.

This is the first demonstration of lymphokines, IFN-gamma and Il2 at the site of a localized autoimmune disease. These observations may indicate that IFN-gamma induced MHC class II antigen expression may serve as a local amplification system in autoimmune and inflammatory eye disease. A better understanding of the role of lymphokines in the mechanisms involved in the development of autoimmunity and inflammation may be beneficial in the treatment of these diseases.

PROJECT NUMBER

Z01 EY 00233-01 LI

PERIOD COVERE	D				
	, 1985 to Sept				
	CT (80 characters or less		· ·		
			of the Retinal Pigment E		
PRINCIPAL INVE			the Principal Investigator) (Name, title labora		
PI:	John J. Hooks	Ph.D.	Head, Section on Immuno.	logy LI, NEI	
			and Virology		
Others;	Barbara Detri	ck Ph.D.	Expert	LI, NEI	
-	Caroline Perc	opo A.B.	Biologist	LI, NEI	
	Yotanna Dalav	anga M.D.	Visiting Fellow	LI, NEI	
	Chi-Chao Chan	M.D.	Staff Ophthalmologist	LI, NEI	
	Garth Stevens	, Jr. M.D.	Senior Staff Fellow	LI, NEI	
COOPERATING L	INITS (d any)				
National :	Institute of D	ental Research	n, Clinical Immunology Se	ction (Siraganian	
Reuben, M					
LAB/BRANCH					
Laborator	y of Immunolog	у			
SECTION					
Section of	n Immunology a	nd Virology			
INSTITUTE AND L	LOCATION				
NEI, NIH,	Bethesda, Mar	yland 20892			
TOTAL MAN-YEAR	RS	PROFESSIONAL	OTHER		
1.4		1.1		0.3	
CHECK APPROPE	RIATE BOX(ES)				
(a) Huma	an subjects	💹 (b) Human tis:	sues 🗌 (c) Neither		
☐ (a1)	Minors				
(a2)	Interviews				
SUMMARY OF WORK (Use standard unreduced type. Do not exceed the space provided.)					

The retinal pigment epithelial (rpe) cell is a major regulatory cell in the eye. That is, the rpe cell exerts a variety of actions in maintaining retinal integrity and function. In order to more effectively study this cell in vivo and in vitro, we have produced monoclonal antibodies directed against human rpe cells.

Using immunoperoxidase assays (ABC), we have identified two mouse IgG monoclonal antibodies which react with the human rpe cell. The monoclonal antibodies are both specific for the rpe cell within the eye, since they do not react with any other ocular structures. Moreover, these antibodies do not cross react with human skin, kidney or peripheral mononuclear cells.

This is the first monoclonal antibody which is directed solely at the human rpe cell. Further characterization and studies with this antibody should prove useful in the identification of rpe cells in situ and in vitro. Moreover, this immunoglobulin will allow us to probe the bioregulatory functions of the cell.

PROJECT NUMBER

Z01 EY 00234-01 LI

PERIOD COVERED	D							
October l	l, 1985 to Sep	tember 30,	1986					
TITLE OF PROJEC	CT (80 charecters or less	Title must fit on o	ne line between	the borders)			
MHC Class	s II Antigens	in the Pat	hogenesi	s of I	nflammatory Di	seases		
PRINCIPAL INVES	TIGATOR (List other pro	fessional personnel	below the Prince	ipal Investig	ator) (Name title, laboral	tory end institute afti	liation)	
PI:	John J. Hook	KS .	Ph.D.		Section on Individual Section on Individual Section on Indiana.	mmunology	Ll,	NE1
Others:	Barbara Deti	rick	Ph.D.	Expe	rt		LI,	NEI
•	Yotanna Dala	ivanga	M.D.	Visi	ing Fellow		LI,	NEI
	Chi-Chao Cha	n	M.D.	Staf:	Ophthalmolog	ist	LI,	NEI
	Robert B. Nu	ssenblatt	M.D.	Depu	y Clinical Di	rector		NE1
LAB/BRANCH	School of Med	licine, loa	nnina, G	reece	(Haralampos M.	noulsopoul	os, M.I	, ,
Laborator	y of Immunolo	gy						
SECTION								
Section o	on Immunology	and Virolo	gy					
INSTITUTE AND LO								
NEI, NIH,	Bethesda, Ma	ryland 20	892					
TOTAL MAN-YEAR	•	PROFESSIONAL			OTHER	_		
0.	56	0.	56			0		ļ
CHECK APPROPRI								
🗌 (a) Huma	n subjects	(b) Huma	in tissues	\propto	(c) Neither			
🔲 (a1) M								
☐ (a2) Ir	nterviews							
SUMMARY OF MO	PK ///se standard ward	was time On ant						

MHC class II antigens, HLA-DR in the human and Ia in the mouse, are membrane bound glycoproteins that are encoded by genes of the major histocompatibility complex. Expression of these antigens is of great functional importance for the initiation and perpetuation of immune responses. In a number of immunopathologic conditions HLA-DR antigen negative cells are stimulated to express class II antigens. In these cases an immunologic role has been postulated for the class II antigen expression.

During the past year, we have determined if class II antigens are expressed in certain diseases and we have evaluated their possible role in autoimmune and inflammatory diseases. Initial studies identified cells in the anterior segment and cells in the retina (rpe cell) which express class II antigens during inflammatory eye diseases. These studies have been extended to evaluate Sjogren's syndrome. We found that the salivary gland in Sjogren's syndrome is infiltrated predominantly by T-lymphocytes and that this is associated with class II antigen expression on glandular epithelial cells.

These studies on MHC class II antigen expression in localized autoimmune diseases provide evidence that the activation of these antigens may contribute to the immunopathogenesis of these disease.

PROJECT NUMBER

Z01 EY 00184-04 LI

PERIOD COVERE				
October	1, 1985 to Septem	ber 30, 1986		
	CT (80 characters or less. Title ri		een the borders)	
Cellular	Mechanisms in Uv	eitis		
PRINCIPAL INVES	STIGATOR (List other profession	el personnel below the P	rincipal Investigator) (Name, title, laboratory, and ins	ititute affiliation)
PI:	Rachel Caspi	Ph.D.	Visiting Fellow	LI, NEI
Others:	Robert B. Nussen	blatt M.D.	Deputy Clinical Director	NEI
	Francois Roberge	M.D.	Visiting Associate	LI, NEI
2	Chi-Chao Chan	M.D.	Staff Ophthalmologist	LI, NEI
	William Leake	M.S.	Biologist	LI, NEI
	Susan Lightman	M.D.	Visiting Fellow	LI, NEI
	Myung Kim	M.D.	Visiting Fellow	LI, NEI
AB/BRANCH	<i>5 7</i>			
Laborato	ry of Immunology			
	on Immunoregulati			
NSTITUTE AND L		011		
	l, Bethesda, Maryl	and 20892		
TOTAL MAN-YEAR	RS PROF	ESSIONAL	OTHER	
1.0)2	1.02	0	
CHECK APPROPE (a) Huma (a1) (a2)	an subjects 🔲 (t	D) Human tissue:	s 🗔 (c) Neither	

SUMMARY OF WORK (Use standard unreduced type. Do not exceed the space provided.)

In vivo functional long-term T-cell lines and T-cell clones are developed and maintained in vitro from both peripheral blood and ocular fluids of humans and animals. The phenotype and functional properties of these cells, as well as their interaction with ocular resident cells are being studied. The goal of these studies will be to identify the immunoreactive cells and mediators involved in the intraocular inflammatory process.

PROJECT NUMBER

Z01 EY 00222-01 LI

PERIOD COVERED						
October 1, 1985 to Sep	tember 30, 19	986				
TITLE OF PROJECT (80 characters or less			borders)			
Kinetics of T-lymphocy	tes in the Ev	es with	Exper	imental Auto	aimmune Uno:	itia
PRINCIPAL INVESTIGATOR (List other pro-	fessional personnel belo	w the Principa	Investigato	r.) (Name, title, fabor	story, and institute at	filiation)
PI: Chi-Chao Chan		M.D.	Staff	Ophthalmol	ogist	LI, NEI
Others: Igal Gery		Ph.D.		Section on inology	Experimenta	al LI, NEI
: Robert B. Nus	senblatt	M.D.	Deputy	Clinical 1	Director	NEI
COOPERATING UNITS (# any)						ĺ
University of Tokyo, S	chool of Medi	cine (Ma	anabu !	lochizuki. N	1.D.): Hadas	ssab Hebrew
University Hospital, D	epartment of	Ophthalr	mology	(David Benl	Ezra, M.D.)	, sour meanew
LAB/BRANCH						
Laboratory of Immunolog	gv					
SECTION		 				
Section on Immunoregula	ation					
INSTITUTE AND LOCATION						
NEI, NIH, Bethesda, Mar	rvland 20892					
TOTAL MAN-YEARS	PROFESSIONAL.		110	IER		
0.07	0.07				0	
CHECK APPROPRIATE BOX(ES)						
(a) Human subjects	(b) Human ti	ssues	□3 (c)	Neither		
(a1) Minors						
(a2) Interviews						ľ
SUMMARY OF WORK (Use standard unred	uced type. Do not excee	d the space p	provided)	· -		
						,

Identity and topographic localization of immunocompetent cells in rats with experimental autoimmune uveoretinitis were analyzed by immunohistochemical studies. The lymphocyte population at the inflammatory sites was found to change markedly during the course of disease. In the early stage, T-helper/inducers are the predominant cells in the eye. A relative increase of T-suppressor/cytotoxic cells in the late stage were observed. These kinetics can be influenced by cyclosporine and dexamethosone treatment.

PROJECT NUMBER DEPARTMENT OF HEALTH AND HUMAN SERVICES - PUBLIC HEALTH SERVICE Z01 EY 00223-01 LI NOTICE OF INTRAMURAL RESEARCH PROJECT 'ERIOD COVERED October 1, 1985 to September 30, 1986 TILE OF PROJECT (80 characters or less. Title must fit on one line between the borders.) Expression of Class II Antigens in Ocular Inflammatory Disorders PRINCIPAL INVESTIGATOR (List other professional personnel below the Principal Investigator) (Name, btle, laboratory, and institute affiliation) PI: Chi-Chao Chan M.D. Staff Ophthalmologist LI, NEI Barbara Detrick Others: Ph.D. Expert LI, NEI John J. Hooks Ph.D. Head, Section on Immunology LI, NEI and Virology Leslie S. Fujikawa M.D. Senior Staff Fellow LI, NEI Igal Gerv Ph.D. Head, Section on Experimental LI, NEI Immunology COOPERATING UNITS (# any) LAB/BRANCH Laboratory of Immunology SECTION Section on Immunoregulation INSTITUTE AND LOCATION NEI, NIH, Bethesda, Marvland TOTAL MAN-YEARS PROFESSIONAL OTHER 0.3 0.3 0

SUMMARY OF WORK (Use standard unreduced type. Do not exceed the space provided.)

(b) Human tissues

CHECK APPROPRIATE BOX(ES)

(a) Human subjects

(a1) Minors (a2) Interviews

Expression of class II antigens on ocular nonlymphoid cells was evaluated in experimental autoimmune uveoretinitis (EAU) by the immunoperoxidase technique. Class II antigens on nonlymphoid cells were not detected from normal rats. However, these antigens were detected on the retinal pigment epithelia and ciliary body epithelia a few days prior to the development of clinical and histopathological EAU. Ia antigen was noted on the retinal vascular endothelia at the onset of cellular infiltration in the retina, and appeared on the corneal keratocytes and scleral fibroblasts after the early stage of clinical EAU.

(c) Neither

The study demonstrates that during the course of EAU the ocular nonlymphoid cells can be activated to express class II antigens. This antigen expression may be important in the initiation and perpetuation of immune reactivity in the eye.

PROJECT NUMBER

Z01 EY 00224-01 LI

October	1, 1985 to September	30, 1986		
TITLE OF PROJE Sympathe	CT (80 characters or less Title must in etic Ophthalmia: Immu	on one line between nopatholo	pen the borders) gical Findings	
PRINCIPAL INVE	STIGATOR (List other professional perso	nnel below the P	rincipal Investigator.) (Name, title, laboratory, and institute	affiliation)
PI:	Chi-Chao Chan	M.D.	Staff Ophthalmologist	LI, NEI
Others:	Robert B. Nussenblat	t M.D.	Deputy Clinical Director	NEI
	Leslie S. Fujikawa	M.D.	Senior Staff Fellow	LI, NEI
-	Alan G. Palestine	M.D.	Head, Section on Clinical Immunology	LI, NEI
	Garth Stevens, Jr.	M.D.	Senior Staff Fellow	LI, NEI
LAB/BRANCH Laborato	ry of Immunology			
Section	on Immunoregulation			
NEI, NIH	OCATION , Bethesda, Maryland	20892		
TOTAL MAN-YEAR 0.	RS PROFESSION	NAL. 0.37	OTHER 0	
		man tissues	c) Neither	
SUMMARY OF WO	ORK (Use standard unreduced type Do	not exceed the s	pace provided)	

Immunocompetent cells in the ocular tissues from six patients with a clinical diagnosis of sympathetic ophthalmia were examined using the immunohistochemical technique. The choroidal infiltrates were composed primarily of T-lymphocytes. Different amounts of macrophages and 13 lymphocytes were present in each case. A varied spectrum of immunopathological and histopathological findings may occur in clinically diagnosed sympathetic ophthalmia.

PROJECT NUMBER

Z01 EY 00225-01 LI

October	l, 1985 to Septembe	ow 30 1096		
	ECT (80 characters or less Title mu		een the borders)	
	lammatory Complicat			
			Principal Investigator) (Name, title, laboratory, and institu	te affiliation)
PI:	Chi-Chao Chan	M.D.		LI, NEI
Others:	Robert B. Nussenbl Leslie S. Fujikawa		ere de la companya de	NEI LI, NEI
-	Richard P. Wetzig			LI, NEI
	Francois Roberge			LI, KEI
LAB/BRANCH				
Laborato	ry of Immunology			
SECTION				
Section	on Immunoregulation	1		
INSTITUTE AND				
NEI, NIH	, Bethesda, Marylan	d 20892 _		
TOTAL MAN-YEA	RS PROFES	0.19	OTHER 0	
☐ (a1)	an subjects 💎 🖾 (b)	Human tissue	s (c) Neither	
SUMMARY OF W	ORK (Use standard unreduced type	Do not exceed the	space provided.)	•

Complications of post-inflammation in uveitis patients included destruction of photoreceptors, gliosis, choroidal scar, and formations of cyclitic membrane, snowbanking and preretinal membrane. Eyes enucleated from patients with end stages of chronic anterior uveitis (formation of cyclitic membrane), pars planitis (formation of preretinal membrane) were evaluated. Glial cells and proliferating Muller cells were the major components in these membranes.

PERIOD COVERED

PROJECT NUMBER

Z01 EY 00226-01 LI

October 1, 1985 to Sep	otember 30, 1986		
TRILE OF PROJECT (80 characters or less	Title must fit on one line between	the borders)	
Immunopathology of Ocu	lar Onchocerciasis		
PRINCIPAL INVESTIGATOR (List other pro	dessional personnel below the Princ	ipal Investigator) (Name, title, laboratory, and institut	te affiliation)
PI: Chi-Chao Cha	an M.D.	Staff Ophthalmologist	LI, NEI
Others: Robert B. No	ussenblatt M.D.	Deputy Clinical Director	NEI
		ious Diseases, Clinical Para	sitic
Diseases Section (Eric LAB/BRANCH Laboratory of Immunolo			
Section on Immunoregul	ation		
NEI, NIH, Bethesda, Ma	aryland 20892		
TOTAL MAN-YEARS 0.2	PROFESSIONAL 0.2	OTHER	
CHECK APPROPRIATE BOXIES) (a) Human subjects (a1) Minors (a2) Interviews	☑ (b) Human tissues	(c) Neither	
SUMMARY OF WORK (Use standard unred	luced type. Do not exceed the space	e provided)	•

Ocular specimens and sera from 12 patients with onchocerciasis and 10 controls were studied. A mild to moderate chronic inflammatory cellular infiltration was present in the conjunctiva of the onchocerciasis patients. T-lymphocytes were the predominant inflammatory cells with the T-suppressor subset being significantly increased in the onchocerciasis patients when compared to controls. In the onchocerciasis patients, the nonlymphoid cells in the conjunctiva and iris, such as vascular endothelia, pericytes and fibroblasts, showed an increase in expression of class II antigens. The anti-onchocerca Volvulus antibodies in the sera and aqueous humor were significantly higher in the patients compared to the controls. These findings suggest that T-cells are important in the ocular immune response to onchocerca and that expression of class II antigens on nonlymphoid cells and the humoral factors may all play a critical role in ocular onchocerciasis.

PROJECT NUMBER

	NOTIC	E OF INTRAMURA	L RESEAR	CH PROJ	ECT	Z01 EY 00	0216-01 LI
PERIOD COVE		to September 30), 1986				
		racters or less Title must fit ronectin in Wour					
PRINCIPAL IN	VESTIGATOR	(List other professional perso	nnel below the				e affiliation)
PI:	Leslie	S. Fujikawa	M.D.	Senior	Staff Fell	OW	LI, NEI
Others:		B. Nussemblatt Datiles	M.D. M.D.		; Clinical D ing Scientis		NEI CB, NEI
COOPERATING	3 UNITS (d an	(y)					
LAB/BRANCH							
	rv of I	mmunology					
SECTION		noregulation					
INSTITUTE AN	D LOCATION	sda, Maryland 2	20892				
TOTAL MAN-Y	EARS	PROFESSIO	VAL		OTHER		
0	.18		0.18			0	
_ `		ects (b) Hu	ıman tissue	es 🗆	(c) Neither		

SUMMARY OF WORK (Use standard unreduced type. Do not exceed the space provided.)

The role of fibronectin in corneal wound healing was studied both in man and in experimental animals. Studies were carried out in order to characterize the possible effect of fibronectin on facilitating wound healing of the epithelium in rabbits. A randomized masked study will evaluate the effectiveness of fibronectin drops in treating recurrent corneal epithelial defects in patients.

PROJECT NUMBER DEPARTMENT OF HEALTH AND HUMAN SERVICES - PUBLIC HEALTH SERVICE Z01 EY 00092-08 LI NOTICE OF INTRAMURAL RESEARCH PROJECT PERIOD COVERED October 1, 1985 to September 30, 1986 TITLE OF PROJECT (80 characters or less. Title must fit on one line between the borders.) HLA, ABO, and B-cell Alloantigens and Ocular Inflammatory Disease PRINCIPAL INVESTIGATOR (List other professional personnel below the Principal Investigator.) (Name title laboratory and institute affiliation) PI: Robert B. Nussenblatt Deputy Clinical Director M.D. NEI COOPERATING UNITS (d any) LAB/BRANCH Laboratory of Immunology Section on Immunoregulation INSTITUTE AND LOCATION NEI, NIH, Bethesda, Marvland 20892 OTHER TOTAL MAN-YEARS **PROFESSIONAL**

SUMMARY OF WORK (Use standard unreduced type. Do not exceed the space provided.)

0.03

CHECK APPROPRIATE BOX(ES)

(a) Human subjects

(a1) Minors (a2) Interviews

Patients with ocular toxoplasmosis, pars planitis, Behcet's disease, chorioretinitis of unknown origin, are being studied to determine the phenotype frequency of the HLA, ABO, and B-cell alloantigens. Because the B-cell alloantigens or DR antigens are thought to play a role in the immunologic response to antigens, these findings will complement other immune uveitis studies being simultaneously carried out.

(c) Neither

0.03

(b) Human tissues

PROJECT NUMBER

Z01 EY 00075-08 LI

October 1,	1985 to Sept	ember	30, 198	36				
	T (80 charecters or less ctions in Ocu							
PRINCIPAL INVEST						igator)(Name title laboratory and in outy Clinical Directo		NEI
Others:	Alan G. Pale	stine		M.D.		d, Section on Clinic mmunology	cal LI,	NEI
	Chi-Chao Cha	n		M.D.	Sta	ff Ophthalmologist	LI,	NEI
	William Leak	e		M.S.	Bio	logist	LI,	NEI
	Shigeto Hiro	se		M.D.	Vis	iting Fellow		NEI
LAB/BRANCH Laboratory	of Immunolog	y						
Section on	Immunoregula	tion						
NEI, NIH,	OCATION Bethesda, Mar	yland	20892					
TOTAL MAN-YEAR 0.8		PROFESS	O. 2	6		OTHER 0.6		
	n subjects finors nterviews		Human ti			(c) Neither		
CHIMMARY OF WO	DK ///en etandard upma	uced hose	Do not over	ad the coac	o proudor	4 1		

OF WORK (Use standard unreduced type. Do not exceed the space provided.)

In vitro cellular immune functions and lymphocyte subsets are being studied in a masked method in patients with ocular toxoplasmosis, pars planitis, Behcet's disease, geographic choroiditis, and chorioretinitis of unknown origin. Crude ocular antigens, as well as purified uveitogenic soluble antigen (S-antigen) and IRBP of the retina, are being used in a lymphocyte microculture technique to evaluate the presence of cellular immune memory to ocular tissues. In addition, purified antigens from the toxoplasmosis organism are also being tested in this in vitro system. A subgroup of patients with posterior uveitis has been identified as having this immunologic memory. Lymphocyte subsets in the blood and in the eye are being defined in these patients by monoclonal antibodies. These results shed light on the basic mechanisms of uveitis and may be used as a quide for specific immunologic therapy. In a small group of selected patients, chorioretinal biopsies are performed to evaluate the on-going ocular immune response.

DEPARTMENT OF HEALTH AND HUMAN SERVICES - PUBLIC HEALTH SERVICE Z01 EY 00094-08 LI NOTICE OF INTRAMURAL RESEARCH PROJECT PERIOD COVERED October 1, 1985 to September 30, 1986 TITLE OF PROJECT (80 characters or less. Title must fit on one line between the borders.) Immune Mechanisms in Experimental Autoimmune Uveitis PRINCIPAL INVESTIGATOR (List other professional personnel below the Principal Investigator) (Name title laboratory and institute attiliation) PI: Robert B. Nussemblatt M.D. Deputy Clinical Director NEI Others: Igal Gerv Ph.D. Head, Section on Experimental LI, NEI Immunology - Cathy McAllister Ph.D. Extramural Fellow LI, NEI Barbara Vistica B.A. Microbiologist LI, NEI William Leake M.S. Biologist LI, NEI COOPERATING UNITS (d any) LAB/BRANCH Laboratory of Immunology SECTION Section on Immunoregulation INSTITUTE AND LOCATION NEI, NIH, Bethesda, Maryland 20892 TOTAL MAN-YEARS PROFESSIONAL OTHER 0.4 0.75 0.35

PROJECT NUMBER

SUMMARY OF WORK (Use standard unreduced type. Do not exceed the space provided.)

(b) Human tissues

CHECK APPROPRIATE BOX(ES)

(a) Human subjects

(a1) Minors (a2) Interviews

Lewis rats and non-human primates, immunized at a site distant to the eye with the retinal soluble antigen (S-antigen) in complete Freund's adjuvant, develop experimental autoimmune uveitis (EAU). Lymph node cells and peripheral lymphocytes from immunized animals manifested significant cellular immune responses measured by the lymphocyte culturing technique. Cyclosporine, a drug with specific anti-T-activity, has been found to be exceptionally effective in protecting rats with EAU, and suppressor cells potentially play a role in this protective mechanism. As well, the inducer cell T-cell fraction in the lymph node appears to be most susceptible to cyclosporine therapy. Attempts at local immunosuppressive therapy in order to prevent EAU have begun. The use of topical CsA has been used to evaluate its effectiveness in EAU. Additionally, newer cyclosporines, particularly D&G, have been evaluated in this model, with their efficacy compared to that of cyclosporine A. Ciamexone, a drug with immunopotentiating characteristics, has always been utilized in this model.

(c) Neither

- 1

PERIOD COVERED

PROJECT NUMBER

201 00115-06 LI

Uctober	1, 1985 to Septemb	er 30,	1986			
TITLE OF PRO.	JECT (80 characters or less. Title i	nust fit on or	e line between	the borders)		
Cyclospo	rine Therapy in Uv	eitis				
PRINCIPAL INV	ESTIGATOR (List other profession	al personnel	below the Prin	cipal Investigator) (Name title laboratory and i	institute affiliation:	
PI:	Robert B. Nussen	blatt	M.D.	Deputy Clinical Directo	r	NEI
Others:	Alan G. Palestin	е	M.D.	Head, Section on Clinic Immunology	al LI,	NEI
-	Garth Stevens, J	r.	M.D.	Senior Staff Fellow	LI,	NEI
	Leslie S. Fujika	wa	M.D.	Senior Staff Fellow	LI,	NEI
	Francois Roberge		M.D.	Visiting Associate		NEI
	Richard P. Wetzi	g	M.D.	Senior Staff Fellow		NEI
LAB/BRANCH Laborato	ry of Immunology					
SECTION Section	on Immunoregulatio	n	-			
NEI, NIH	LOCATION , Bethesda, Maryla	nd 208	92			
TOTAL MAN-YE	2.52 PROF	ESSIONAL	2.52	OTHER (0	
(a) Hun	PRIATE BOX(ES) man subjects	b) Huma	n tissues	(c) Neither		

SUMMARY OF WORK (Use standard unreduced type. Do not exceed the space provided.)

Cyclosporine, an endecapeptide fungal product with specific anti-T-cell characteristics, will be administered to patients with sight-threatening ocular inflammatory disease of non-infectious origin who have failed on either corticosteroid or cytotoxic agent therapy. This will be done to test cyclosporine's efficacy in the treatment of uveitis. Within the context of these ongoing studies, the effect of hydergine on reversing cyclosporine induced nephrotoxicity is being evaluated in a randomized, masked, cross-over study. Additionally, selected patients whose uveitis is well controlled on cyclosporine for one year or more are undergoing kidney biopsies to evaluate the long term effects of this agent.

		•

PROJECT NUMBER

201 EY 00159-04 LI

PERIOD COVE	RED			
October	1, 1985 to September	er 30, 1986		•
	JECT (80 characters or less. Title n		n the borders)	
Cvclospo	orine Therapy of Chi	ildhood Uveiti	S	
PRINCIPAL IN	ESTIGATOR (List other professions	al personnel below the Pril	ncipal Investigator) (Name title laboratory and institut	te affiliation)
PI:	Robert B. Nussenb	latt M.D.	Deputy Clinical Director	NEI
Others:	Alan G. Palestine	M.D.	Head, Section on Clinical Immunology	LI, NEI
•	Chi-Chao Chan	M.D.	Staff Ophthalmologist	LI, NEI
	William Leake	M.S.	Biologist	LI, NEI
SECTION	ory of Immunology on Immunoregulatio	n		
NEI, NII	D LOCATION I, Bethesda, Maryla	nd 20892		
TOTAL MAN-Y	EARS PROF	0.15	OTHER 0.05	
(a) Hu	man subjects (I) Minors Interviews	b) Human tissues	(c) Neither	
	WQRK (Use standard unreduced to	ype. Do not exceed the sp	pace provided)	
charac	teristics, is being	g tested in ch	epeptide with specific anti-T ildren with sight-threatening origin who have failed on co	ocular

inflammatory disease of non-infectious origin who have failed on therapy.

PROJECT NUMBER Z01 EY 00215-01 LI

October 1	, 1985 to September 30,	1986			
	T (80 characters or less. Title must tit on one d. Double-Masked Study of			dogenous Uvei	itis
PRINCIPAL INVEST	GATOR (List other professional personnel be Robert B. Nussenblatt				NEI
Others:	Alan G. Palestine	M.D.	Head, Section on C Immunology	linical l	LI, NEI
	Garth Stevens, Jr.	M.D.	Senior Staff Fello	w l	LI, NEI
-	Leslie S. Fujikawa	M.D.	Senior Staff Fello	w I	LI, NEI
	Chi-Chao Chan	M.D.	Staff Ophthalmolog	ist l	LI, NEI
	Richard P. Wetzig	M.D.	Senior Staff Fello	w I	LI, NEI
LAB/BRANCH					
	y of Immunology				
SECTION					
Section o	n Immunoregulation				
NEI, NIH,	xation Bethesda, Maryland 208	92			
TOTAL MAN-YEARS		. 6	OTHER	0	
CHECK APPROPRIEMS (a) Human (a1) M	n subjects 🔲 (b) Human	tissues	☐ (c) Neither		

SUMMARY OF WORK (Use standard unreduced type. Do not exceed the space provided.)

Cyclosporine's efficacy in the treatment of severe endogenous uveitis was evaluated in this randomized, double-masked study. The study will evaluate the effectiveness of cyclosporine therapy to that of systemic corticosteroid administration. Patients meeting the entry requirements were randomized to either cyclosporine or corticosteroid therapy. Patients are evaluated at three months in order to determine whether they were therapeutic "successes" or not. If not, the patients are then treated with the alternate medication.

PROJECT NUMBER

Z01 EY 00228-01 LI

PERIOD COVER							
October	1, 1985 to Sep	otember 30,	1986				
TITLE OF PROJE	CT (80 characters or les	s. Title must fit on one	line between ti	he borders)			
Study of	Ocular Glial	Cells Invol	vement in	n Uveit	is		
PRINCIPAL INVE	STIGATOR (List other pri	olessional personnel b	elow the Princip	al Investigati	or) (Name, title, labo	oratory, and institute at	Yeleton)
PI:	Francois Robe	erge	M.D.	Visiti	ng Fellow		LI, NEI
Others:	Robert B. Nus	ssenblatt	M.D.	Deputy	Clinical D	irector	NEI
	Chi-Chao Char	1	M.D.	Staff	Ophthalmolo	gist	LI, NEI
-	Rachel Caspi		Ph.D.	Visiti	ng Fellow		LI, NEI
COOPERATING U	JNITS (d eny)						
Laborator	ry of Immunolo	gy					
Section (on Immunoregul	lation					
NEI, NIH	LOCATION , Bethesda, Ma	aryland 208	92				
TOTAL MAN-YEA	RS	PROFESSIONAL		ОТ	HER		
1	.06	1.0)6			0	
🔲 (a1)	RIATE BOX(ES) an subjects Minors Interviews	(b) Human	tissues	∑ (c) Neither		
SUMMARY OF W	ORK (Use standard unrec	duced type. Do not ex	ceed the space	provided)			•

The work concerns uveoretinal inflammatory disease mechanisms. The study focused on evaluating the role of resident cells at the target organ level. Muller cells interactions with T-lymphocyte cell line was studied in vitro. opposite effects of Muller cells on T-lymphocytes proliferation were found that could be expressed under different culture conditions. An inhibitory effect on antigen driven proliferation of T-helper lymphocytes could be modulated with various drugs that allowed the Muller cells to function as antigen presenting cells.

PROJECT NUMBER

Z01 EY 00227-01 LI

PERIOD COVERE					
October	1, 1985 to September 30,	1986			
	CT (80 characters or less. Title must fit on on				
Histopat	hology of Pars Planitis	and Exp	erimental Autoimmune Uveitis		
			cipal Investigator) (Name, title, laboratory, and institute a	Tiliation)	
PI:	Richard P. Wetzig	M.D.	Senior Staff Fellow	LI, NE	I
Others:		M.D.	Deputy Clinical Director	NE:	I
	Chi-Chao Chan	M.D.	Staff Ophthalmologist	LI, NE	I
		Ph.D.	Expert	LI, NFI	I
	John J. Hooks	Ph.D.	Head, Section on Immunology and Virology	LI, NEI	I
LAB/BRANCH					
	ry of Immunology				
SECTION					
Section o	on Immunoregulation				
NEI, NIH	ocation , Bethesda, Maryland 20	892			
TOTAL MAN-YEAR	/ m	67	OTHER 0		
🔲 (a1) l	MATE BOX(ES) an subjects · & (b) Huma Minors nterviews	n tissues	☐ (c) Neither		

SUMMARY OF WORK (Use standard unreduced type. Do not exceed the space provided.)

Studies in animals and in patients are being carried out to determine factors influencing ocular immune responses. In an animal model, rats are immunized with S-retinal antigen to produce experimental autoimmune uveitis. Animals in one group received anti-Ia antibody intraperitoneally and developed the onset of uveitis significantly later and to a lesser extent than controls. Histopathologically, the anti-Ia treated animals had much less inflammation than did controls. A human eye with pars planitis was also studied immunohistologically. In the pars plana region there was an elevated helper to suppressor T-cell ratio. In addition, the snowbank area showed staining for glial fibrillary acid protein Muller cells, type IV collagen and laminin. There was staining for HLA-DR throughout the globe. The results of these studies shed light on how surface antigens effect and are transmitted by ocular immune responses.

PROJECT NUMBER

Z01 EY 00201-02 LMOD

PERIOD COVE					
October	1, 1985 to Septe	ember 30, 1986			
TITLE OF PRO	DJECT (80 characters or less	Title must fit on one line	between the borde	ers)	
Molecular	r Biology of Alo	dose Reductase			
PRINCIPAL IN	VESTIGATOR (List other pro	fessional personnel below	the Principal Inves	stigato") (Name title laboratory and in	strute affiliation)
PI:	Deborah Carper		Ph.D.	Biologist	LMOD, NEI
				•	
Others:	Toshimichi Shir	nohara	Ph.D.	Biologist	LMDB, NEI
-	Cheryl Craft		Ph.D.	Guest Worker	MIDP, NICHHD
	Jin H. Kinoshi	ta	Ph.D.	Scientific Director	NEI
ļ					
COOPERATING	S UNITS (d any)				
None					
LAB/BRANCH					
Laborato	ry of Mechanism	s of Ocular Di	sease		
SECTION	-				
Section	on Cataracts				
INSTITUTE AN	D LOCATION	·			
NEI, NIH	, Bethesda, Mar	yland 20892			
TOTAL MAN-Y		PROFESSIONAL		OTHER	
	1.2	1.2		0	
	OPRIATE BOX(ES)				
	man subjects	(b) Human tis	sues 🗔	(c) Neither	
🔲 🔲 (a1) Minors				
│ □ (a2) Interviews				
SUMMARY OF	WORK (Use standard unred	uced type. Do not exceed	the space provide	ed)	

Aldose reductase (AR), an enzyme which converts sugars to sugar alcohols, has been implicated in diabetic complications of the eye and the peripheral nerves. Our studies have focused on isolating and characterizing the gene for aldose reductase. As an initial step, putative AR cDNA clones from a bovine retina cDNA kgt 11 library were isolated and sequenced. Sequencing of three non-homologous inserts yielded approximately 700 nucleotides or around one-third of the full length mRNA. Northern analysis showed that the full size of AR mRNA is around 2 kilobases in the bovine retina and slightly larger in the bovine brain. Homologies between the bovine retina AR cDNA and mRNAs from cultured dog lens epithelial cells and cultured rabbit kidney cells have not been detected. Restriction enzyme analysis of total bovine genomic DNA indicates that fewer than 3 genes are present.

PROJECT NUMBER

DEPARTM	ENT OF HEALTH AND HO	MAN SERVICES - PUBL	IC HEALTH SERVICE	
	NOTICE OF INTRAMU	JRAL RESEARCH P	PROJECT	EY 00236-01 LMOD
PERIOD COVERED				
	1985 to September			
TITLE OF PROJECT	(80 characters or less. Title mu	ist fit on one line between th	e borders)	
Philly Mous				
PRINCIPAL INVEST	IGATOR (List other professional	personne' below the Princip.	al Investigator) (Name title laboratory	and institute affiliation
PI:	Deborah Carper	Ph.D.	Biologist	LMOD, NEI
Others:	George Inana	M.D. Ph.D.	Medical Officer	LMOD. NET
-			Scientific Director	
COOPERATING UN	ITS (r any)			
John Clark	Ph.D., faculty Un	iversity of Was	hington School of Med	icine
Mike Gorin	M.D., Ph.D. UCLA	School of Medic	ine	
LAB/BRANCH				
	of Mechanisms of	<u>Ocular Disease</u>		
SECTION				
Section on		·		
INSTITUTE AND LO				
	ethesda, Maryland			
TOTAL MAN-YEARS		SSIONAL	OTHER	
0.		0.2		0
CHECK APPROPRIA	_			
(a) Human	r subjects 🗀 (b)	Human tissues	😠 (c) Neither	

SUMMARY OF WORK (Use standard unreduced type. Do not exceed the space provided.)

☐ (a1) Minors (a2) Interviews

Studies are continuing on the Philly mouse which develops a dominantly inherited cataract. The cataract is visible by 35 days of age. Microscopic changes in the lens occur much earlier and include failure of lens fiber cells to elongate. We have been investigating several facets of cataract development in the Philly mouse including the composition of crystallin proteins at different stages in lens growth, synthesis and translational efficiencies of lens mRNAs, and phase transition temperatures in the Philly mouse and F1 heterozygote.

2-D gel electrophoresis of Philly mouse lens proteins indicate a specific deficiency of a 27K basic 2-crystallin and its functional mRNA. Interestingly, a unique more acidic 26K protein and its functional mRNA is present. The relationship between these two proteins is now being investigated using a zB_n cDNA probe.

The behavior of the lens cytoplasm phase transition temperature (Tc) was different in the Swiss-Webster, Philly, and Swiss Webster x Philly heterozygotes. The slope (dTc/dt) changes from negative to positive on day 27 for the Philly and day 38 for the hybrid, just prior to cataract in these animals. The slope continues to be negative in the Swiss Webster. The change in slope is a graphical confirmation of the disturbance in lens cell composition and an early indicator of conditions which lead to opacity.

PROJECT NUMBER

ZO1 EY 00189-03 LMOD

PERIOD COVERED					
October 1, 1985 to Sept	ember 30, 1986				
TITLE OF PROJECT (80 characters or les					
Protein Kinases in Lens	Function and 0:	xidation of	Proteins	in Cataractogenes	318
PRINCIPAL INVESTIGATOR (List other pri	ofessional personnel below th	he Principal Investigi	ator) (Name title la	aboratory and institute affiliatio	n)
- 1	D = D	Eveent	TMOD	MET	
Donita L. Garland	Ph.D	Expert	LMOD,	NEI	
			•		
•					
COOPERATING UNITS (if any)					
None					
LAB/BRANCH	e of Coular Die	0350			
Laboratory of Mechanism	S of Ocular Dis				
SECTION					
Section on Cataracts					
INSTITUTE AND LOCATION					
NEI, NIH, Bethesda, Mar	yland 20892				
TOTAL MAN-YEARS	PROFESSIONAL		THER	_	
1.0	1.0			0	
CHECK APPROPRIATE BOX(ES)	<u> </u>				
(a) Human subjects	💢 (b) Human tiss	ues 🗌 ((c) Neither		
(a1) Minors					
(a2) Interviews					
SUMMARY OF WORK (Use standard unre	duced type. Do not exceed t	the space provided)			
The male of enotein kin	* *			s is heing	

The role of protein kinases in regulating metabolism in lens is being addressed by studying: 1) the protein kinases, and 2) the endogenous proteins that serve as substrates for the lens protein kinases. The focus of this study is on the purification of the protein kinases and the characterization of the phosphorylation of four endogenous substrates by cAMP-dependent protein kinases. The phosphorylated proteins are α -crystallin and 26K and 19K intrinsic membrane proteins. Comparison of the amino acid compositions of the 26K and 19K proteins suggest they are closely related. Detailed structural studies are in progress to determine how similar they are. Compounds that are thought to regulate the function of the 26K protein in vivo modulate the phosphorylation of the protein in vitro.

Oxidative changes of lens proteins are thought to occur with aging and to contribute to the development of cataracts. The goals of this project are to determine: 1) the extent of oxidative modification of crystallins and metabolic enzymes in both normal and cataractous lenses; 2) the nature of the modifications and mechanisms leading to the changes; 3) the effect of the modifications on structure and function of lens protein. Bovine and human lenses were used. Incubation of crystallins with ascorbate - FeCl3-02 caused nondisulfide crosslinking of α and β and the partial degradation of all 3 crystallin fractions. Conversion to more acidic species occurred with all 3 fractions, and nontryptophan fluorescence was produced in $\beta_{\rm H}$ fraction. Longer incubations of homogenate with ascorbate - Fe-02 mimicked changes similar to those in brunescent lenses. Proteins became brownish, insoluble and there was an increased carbonyl content.

PROJECT NUMBER

Z01 EY 00237-01 LMOD

PERIOD COV October	VERED 1, 1985 to Septe	mber 30, 1986		
TITLE OF PR Charact	ROJECT (80 characters or less erization of the	Title must fit on one line between Primate Lens	en the borders)	
PRINCIPAL I	NVESTIGATOR (List other profe	ssional personnel below the Pri	incipal Investigator) (Name title, laboratory and	mstitute affiliation
PI:	Paul Russell	Ph.D.	Research Chemist	LMOD, NEI
	-			
COOPERATI	NG UNITS (if any)			
LAB/BRANCH Laborat	ory of Mechanisms	of Ocular Diseas	ses	
SECTION Section	on Cataracts			
NEI, NI	ND LOCATION H, Bethesda, Mary	land 20892		
TOTAL MAN-	YEARS 1.0	PROFESSIONAL 1.0	OTHER 0.0	
☐ (a) H	nopriate Box(Es) uman subjects 1) Minors 2) Interviews	(b) Human tissues	(c) Neither	

SUMMARY OF WORK (Use standard unreduced type. Do not exceed the space provided.)

Developmental differences in the composition of lens proteins have been observed with both monkey and human lenses. These alterations in composition were similar to those observed in rodent lenses at similar stages in development. In order to investigate the role in which these changes may affect optical clarity, the lens crystallin proteins as well as the glycoproteins from the membrane of the lens cells have been studied. Differences in the crystallin composition may give an indication of factors involved in congenital cataracts. Changes in individual crystallins may also be important to understand the aging process in the lens.

 β - and γ - crystallin compositions are changing in the early embryonic period of the primate lens. The β -crystallins have much higher apparent molecular weights in the fetal lens perhaps indicative of a different organization of the subunits of these proteins during this stage of development. In addition, the main γ -crystallin in the embryonic lens is only a very minor component in the adult lens. As the lens cells differentiate into fiber cells, the glycoprotein composition on their membranes changes. Specific glycoprotein differences have been seen in cataractous rodent lenses compared with normal lens; and using lectins, primate lenses can now be probed for changes related to cataract formation and development.

PROJECT NUMBER
Z01 EY 00105-07 LMOD

Sept Sept	ember 30, 1986					
TITLE OF PROJECT (80 characters or less Structure and Compositi	s Title must fit on one line of on of Lens Crys	allins w	ith Respec	t to Catara	ctogene:	sis
PRINCIPAL INVESTIGATOR (List other pr PI: J. Samuel Zig	ofessional personnel below to ler, Jr. Ph.I	e Pnncipal Inves). Res	tigator)(Name tite search Bio	e laborator, and in: logist	stitute affiliatio LMOD,	NEI
Others: Valerie A. Lu _ Qing-ling Hua			siting Fell siting Fel		LMOD, LMOD,	
COOPERATING UNITS (if any) Jules Stein Eye Institu Adelphi University (F. Tennessee (H.M. Jerniga LAB/BRANCH Laboratory of Mechanism	Bettelheim); Dep	partment (
Section on Cataract						
NEI, NIH, Bethesda, Mar	yland 20892					
TOTAL MAN-YEARS 2.8	PROFESSIONAL 2	. 8	OTHER	0.0		
CHECK APPROPRIATE BOX(ES) (a) Human subjects (a1) Minors (a2) Interviews	☑ (b) Human tiss	ues 🗆	(c) Neither			
SUMMARY OF WORK (Use standard unre The ocular lens normall is manifested in the oc during aging and partic characterizing such pro them. Model systems ar or in crystallin soluti activated oxygen are in very probably in the le activated oxygen specie ie, inside the lens or among the oxidant speci therapeutic means to pr	y exists in an ocurrence of markularly during catein changes and e used to study ons. Complex involved in the dans in vivo as we so produce dar in the external es and lens compevent oxidative	environment de de lucidate changes interaction amage procedu. The mage dependents is damage.	nt of high tive modif evelopment ting the m produced in ns among ti duced in potential nds upon wi Understands essential	ication to . We are e echanisms w n organ cul he various such model of the var here they a ding the in l if we are	lens prompted and the prompted to the prompted	oteins in oduce enses f and rated, ons ise
The study of animal colinsights into processes nuclear cataracts prese Huang has isolated and they can be compared widiscovery of a new crys possibilities for study	of cataractoger nt congenitally described the cr th crystallins : tallin not pres	nesis. We in a colorystalling from the cent in ot	e have beg ony of guin s from norm cataractou her specie:	un to inves nea pigs. mal guinea s animals.	tigate Dr. Q-L pigs so The	the •

Transport processes are vital to the maintenance of normal lens homeostasis. Dr. V.A. Lucas is studying the membrane protein present in lens which transports glucose across cell membranes. This glucose transporter has been isolated and an antibody raised against it. Characterization of glucose transport in monkey lens membranes has been done using specific binding assays and known inhibitors of the glucose transporter.



PROJECT NUMBER

Z01 EY 00193-03 LMOD

PERIOD COVERE	D					
October 1	, 1985 to Sept	ember 30, 1	986			
TITLE OF PROJEC	CT (80 characters or less	Title must fit on one	line between the borde	rs)		
Molecular	Biology of He	reditary Ey	e Diseases			
PRINCIPAL INVES	TIGATOR (List other pro	fessional personnel be	olow the Principal Inves	rigator) (Name title, laboratory and in	istitute affiliationi	
PI:	George Inana		M.D., Ph.D.	Section Head	LMOD,	NEI
Others:	Seiichi Totsu	ıka	M.D., Ph.D.	Visiting Fellow	LMOD,	NEI
•	Carmelann Zin	itz	Ph.D.	Staff Fellow	LMOD,	NEI
-	Yoshihiro Hot	ta	M.D.	Visiting Fellow	LMOD,	NEI
	T. Michael Re	dmond	Ph.D.	Staff Fellow	LRCMB,	NEI
See next p	oage.					
LAB/BRANCH						
Laboratory	of Mechanism	s of Ocular	Diseases			
SECTION						
Molecular	Pathology Sec	tion				
INSTITUTE AND L	OCATION					
NEI, NIH,	Bethesda, MD	20892				
TOTAL MAN-YEAR	RS	PROFESSIONAL		OTHER		
	3.6		3.6	0		
CHECK APPROPE	RIATE BOX(ES)					
(a) Huma		X (b) Human	tissues \square	(c) Neither		
☐ (a1) !	Minors					
☐ (a2) I	nterviews					
SUMMARY OF WO	DRK (Use standard unred	luced type. Do not ax	ceed the space provide	d)		

Ornithine Aminotransferase Deficiency in Gyrate Atrophy: Gyrate atrophy (GA) is an autosomal recessive degenerative disease of the retina and choroid of the eye which leads to blindness. We have isolated a gene probe for the human ornithine aminotransferase (OAT), a mitochondrial enzyme which is deficient in GA patients. The gene probe is a \gt11 cDNA clone which was obtained from our human cDNA library through a Western screening method using the anti-human OAT antibodies. The OAT cDNA is 2073 base pairs long and contains the complete coding sequence of the protein. The cDNA-derived OAT sequence is a precursor containing a leader sequence like other mitochondrial enzymes, matches the sequences of seven purified tryptic peptides of pure OAT, and shows homology with another mitochondrial enzyme, aspartate aminotransferase. Examination of the genomic organization of OAT using the cDNA as a probe revealed a gene family consisting of approximately four copies of OAT or OAT-like gene sequences. The OAT gene sequences were mapped to multiple chromosomal loci, confirming the presence of a gene family. Examination of the OAT genes of GA patients has revealed restriction fragment length polymorphisms but no grossly obvious abnormalities, including deletions. Characterizations of multiple gene clones of OAT and the status of OAT mRNA synthesis in GA patients are in progress.

Hereditary Retinoblastoma: We are investigating the molecular basis of malignant transformation in hereditary retinoblastoma using cell culture and molecular genetic techniques. To determine if retinoblastoma has a dominant or recessive malignant phenotype, retinoblastoma cells (Y79) were fused with non-malignant cells (NIH3T3), and the growth characteristics of the hybrid cells were studied. The hybrid cells, containing both the neomycin and GPT markers from the parents, are anchorage-dependent, have a fibroblastic morphology and do not grow in soft agar like the non-malignant parent. The results indicate the malignant phenotype of

retinoblastoma to be recessive.

PROJECT NUMBER

Z01 EY 00003-14 LMOD

PERIOD COVERED October 1, 1985 to September 30, 1986
TITLE OF PROJECT (80 characters or less. Title must fit on one line between the borders.)
Pharmacology of Ocular Complications
PRINCIPAL INVESTIGATOR (List other professional personnel below the Principal Investigator.) (Name: title: laboratory, and institute affiliation.)
PI: Peter F. Kador Ph.D. Research Chemist LMOD, NEI
COOPERATING UNITS (if any) None
LAB/BRANCH Laboratory of Mechanisms of Ocular Diseases
Section on Molecular Pharmacology
INSTITUTE AND LOCATION NEI, NIH, Bethesda, Maryland 20892
TOTAL MAN-YEARS 3.3 PROFESSIONAL 2.8 OTHER 0.5
CHECK APPROPRIATE BOX(ES) (a) Human subjects (b) Human tissues (c) Neither (a1) Minors (a2) Interviews
SUMMARY OF WORK (Use standard unreduced type Do not exceed the space provided) Studies are being conducted on events leading to the onset of various ocular diseases and on methods for their potential pharmacological control. The relationships between the enzymes aldose reductase and aldehyde reductase and the progression of ocular complications such as retinopathy, cataract, pupil function and iris changes, and keratopathy induced by diabetes or galactosemia are being investigated. Methods for either delaying or preventing the onset of these

complications through the pharmacological control of aldose reductase are also

Events leading to the formation of several types of cataracts are also being studied as well as methods for controlling the onset of these cataracts through

being developed.

pharmacological intervention.

PERIOD COVERED

PROJECT NUMBER

Z01 EY 00149-13 LMOD

October 1	, 1985 to Sept	tember 30, 1	986				
	ECT (80 characters or les octure and Fund				s) ssues of the Eye		
PRINCIPAL INVE	ESTIGATOR (List other pr	rofessional personnel b	elos the Princ	ipal Investi	gator) (Name title laboratory and institute	affiliation)	
PI:	W. Gerald Rol	bison, Jr.	Ph.D.		Chief, Section on Pathophysiology	LMOD,	NEI
Others:	Martin L. Ka Masao Nagata Thomas C. Hol		Ph.D. M.D., F Ph.D.	h.D.	Staff Fellow Visiting Associate Senior Staff Fellow		NEI
COOPERATING	UNITS (đ any)						
LAB/BRANCH							
Laborator	y of Mechanism	ms of Ocular	Disease	s			
Section o	n Pathophysio	logy					
NEI, NIH,	LOCATION Bethesda, Mar	ryland 2089	2				
TOTAL MAN-YEA	5.2	PROFESSIONAL	5.0		OTHER .2		
(a1)	PRIATE BOX(ES) nan subjects Minors Interviews	⊠ (b) Humar	tissues		(c) Neither		
SUMMARY OF V	VORK (Use standard unre	duced type. Do not as	cood the space	e provided)		

Aldose reductase has been implicated in two histopathological hallmarks of diabetic retinopathy involving retinal capillary walls: 1) the selective loss in numbers of mural cells (intramural pericytes) from the capillaries; and 2) the thickening of the basement membranes which envelop the cells of the capillary walls. Mural cells contain aldose reductase, accumulate sorbitol, and appear to be more susceptible to incubation in high glucose than are endothelial cells. A thickening of capillary basement membrane ultrastructurally similar to that characteristic of diabetic retinopathy was induced in rat retinas by galactose feeding and was prevented by two structurally different inhibitors of aldose reductase. The diabetic-like thickening of retinal capillary basement membranes in galactose-fed rats was accompanied by other ultrastructural alterations mimicking changes typical of diabetic microangiopathy, such as multilamination, banding of collagen, and the formation of vacuoles and dense inclusions. Bovine, canine, and human mural cells and endothelial cells from retinal capillaries have been grown in cell culture so that the role of aldose reductase in basement membrane synthesis and in various complications of the diabetic state could be studied under chemically defined conditions. Aldose reductase inhibitors are useful for studies of the possible prevention of diabetic retinopathy by oral drugs.

ANNUAL REPORT NATIONAL EYE INSTITUTE October 1, 1985 - September 30, 1986

REPORT OF THE CHIEF, LABORATORY OF MOLECULAR AND DEVELOPMENTAL BIOLOGY
Joram Piatigorsky, Ph.D.

This is the fifth year for the Laboratory of Molecular and Developmental Biology. The three Sections (established in FY 1986) have established autonomy while remaining integrated in their life-styles and their common efforts to study the visual system at the molecular level. The Thursday morning doughnut-data sessions have continued to keep everyone abreast of on-going work and use the collective knowledge of the laboratory members. The invited seminars on alternate Tuesdays was organized this year by Dr. Teresa Borras and featured a stimulating series of talks, while our journal club on the other Tuesdays continued to keep us informed on recent developments in this rapidly moving area of science.

Each Section made notable gains in their research this year, as indicated in the individual reports given in this volume. Some of the recent progress includes not just new findings but reflects a growing clarity in direction. For example, while my group continues to characterize the structure and organization of crystallin genes—ie, to clone and sequence—we have advanced our efforts to include function. Crystallin gene promoters are being studied simultaneously in cell—free systems, in cultured cells and in transgenic mice. Each approach has its advantages and pitfalls. By combining the methods we hope to active a deeper understanding of the regulated, tissue—specific nature of crystallin gene expression. Acquisition of an oligonucleotide synthesizer has also had an impact on our work in that we can now make mutations in cloned gene sequences more easily and cheaply. Despite the considerable expenses of this equipment, it has already paid for itself and been of use to several laboratories in the NEI and other institutes.

In addition to continuing to study the metabolic control of lens cell growth and differentiation, Dr. Peggy Zelenka's section has been increasingly incorporating molecular genetics in their work. Their initial finding reported last year that the oncogene mRNA c-myc is transiently increased during lens fiber cell differentiation has been developed and expanded to other oncogenes. They have found that several oncogenes are expressed in the developing lens, each with a characteristic pattern during differentiation. The role of the oncogene products in the lens is not known yet, but the combined approach of molecular and biochemical investigations should yield new insights to the regulation of lens cell growth and differentiation.

Last year Dr. Toshimichi Shinohara's group reported their initial studies on S-antigen of the retina. They identified several cDNA clones as S-antigen and begun in situ hybridization experiments on the localization of opsin and S-antigen mRNAs in the retina and pineal gland. This year Dr. Shinohara's group has obtained considerably more sequence information on S-antigen and isolated S-antigen genes from the mouse and human, opening new possibilities for the study of tissue-specific gene expression in the retina. Growing knowledge of the sequence of S-antigen has permitted the synthesis of peptides

of S-antigen. These, in turn, are being used to define sites of S-antigen which are responsible for the generation of uveitis in experimental rats. Thus, Dr. Shinohara's basic investigations are being productively coupled with studies having direct relevance to medicine.

In FY 1985 we began a very fruitful collaboration with Dr. Heiner Westphal and his colleagues in the Laboratory of Molecular Genetics (NICHD) for testing crystallin promoters in transgenic mice. These studies have now demonstrated the extreme tissue-specificity of the murine αA -crystallin promoter and its ability to regulate foreign genes (bacterial and viral) in transgenic mice. In one case, transgenic mice were made which contained the aA-crystallin promoter fused to the T-antigen gene of SV40. The lenses of these mice were neoplastically transformed, showing for the first time that the lens is not refractive to cancer, as was long believed. More importantly, the transgenic mice experiments established that crystallin promoters can be used to modify lens genotype and phenotype by genetic engineering. In addition, our results suggest strongly that regulatory sequences for other genes, for example S-antigen or opsin which are expressed in the retina, can also be used to direct foreign genes to different ocular tissues. Thus, it has become possible to consider genetic engineering for both basic and potentially clinical investigations on the eye. We are in the process of developing a facility to produce transgenic mice in our laboratory. Dr. Ana Chepelinsky has been very much involved in this effort. Hopefully, FY 1987 will witness the birth of many interesting LMDB transgenic mice.

Finally, the ARVO electorate has been kind to us this year. Dr. Peggy Zelenka has been elected to the program committee (1987-1989) of the Lens Section and I have been elected as a trustee (1986-1990) representing the Lens Section. This will give the LMDB an administrative voice contributing to the advancement of vision research. Numerous collaborations with other laboratories (listed in the individual reports) have expanded our scope of research and are much appreciated.

DEDUCE COVERED

PROJECT NUMBER

Z01 EY 00127-10 LMDB

October 1, 1985 to Se	ptember 30, 1986			
TITLE OF PROJECT (80 characters or le			- /	
Plasma Membrane Compo				
PRINCIPAL INVESTIGATOR (List other p				
PI: Peggy Zelen	ka Ph.	D. G	eneticist	LMDB, NEI
Luke Pallan	sch Ph.	D. S	taff Fellow	LMDB, NEI
- Malini Vata	l Ph.	D. V	isiting Fellow	LMDB, NEI
Jodell Boyl	e B.S	S. M	edical Student	HHMI
COOPERATING UNITS (d env)				
Beltsville Agricultur	al Research Cente	er, Betls	ville, MD (A. Ferre	etti)
LAB/BRANCH				
Laboratory of Molecula	ar and Developmen	it Biolog	У	
SECTION				
Section on Cellular D	ifferentiation			
NEI, NIH, Bethesda, M	aryland 20892			
TOTAL MAN-YEARS	PROFESSIONAL		OTHER	
2.3		1.5		0.8
CHECK APPROPRIATE BOX(ES) (a) Human subjects (a1) Minors (a2) Interviews	(b) Human tissu	ies 🗆	(c) Neither	
SUMMARY OF WORK (Use standard uni	educed type. Do not exceed th	e space provide		

This project seeks to determine whether the regulation of lens fiber differentiation and maturation is associated with alterations in the plasma membrane. The composition, biosynthesis, and metabolism of lens lipids have been investigated using embryonic and adult chicken lenses, and cultured lens epithelial cells derived from the Nakano mouse. The rate of degradation of the membrane phospholipid, phosphatidylinositol, has been shown to be tightly coupled to the rate of lens epithelial cell division and to cease when the epithelial cells differentiate to form lens fibers. Cultured lens epithelial cells and cultured fibroblasts have been shown to possess a mechanism for the rapid, transient gradation of phosphatidylinositol, which is independent of phospholipase C or phospholipase A_2 . A similar pathway may play a role in the differentiation of lens epithelial cells into lens fibers. Since phosphatidylinositol is rich in arachidonic acid, a precursor of prostaglandins and leukotrienes, the metabolities of arachidonic acid produced by lens epithelial cells are being characterized in an effort to understand the physiological role of phosphatidylinositol degradation. Analysis of the archadonic acid metabolities of cultured lens epithelial cells of several species revealed the presence of both cyclo-oxygenase and lipoxygenase products, including prostagladins E_2 and $F_{2\alpha}$, and leukotrienes. All lens epithelial cell types examined synthesized products of the 5-lipoxygenase pathway of arachidonic acid metabolism. One product of this pathway, 5-hydroxytetraenoic acid, was weakly mitogenic when added to cultured lens epithelial cells in the absence of serum or growth factors. Alterations in phosphatidylinositol metabolism and in the production of arachidonic acid metabolities are being correlated with the action of growth factors in regulating cell division and differentiation.



PROJECT NUMBER

Z01 EY 00238-01 LMDB

October	ED 1, 1985 to Sep	tember 30,	1986		
Proto-on	ECT (80 characters or less cogene Express	Title must fit on one ion During	line between the Lens Diff	ne borders) 'erentiation and Develo	opment
PRINCIPAL INVE	STIGATOR (List other pro Peggy Zelenk		elow the Princip Ph.D.	ad Investigator) (Name title laborator, a Geneticist	ind institute administron) LMDB, NEI
Others:	Luke Pallans Malini Vatal Pravendra Na		Ph.D. Ph.D. Ph.D.	Staff Fellow Visiting Fellow Visiting Fellow	LMDB, NEI LMDB, NEI LMDB, NEI
COOPERATING O			Ph.D.	Uniformed Services of the Health Sci Bethesda, MD	•
Laborator	ry of Molecula	r and Devel	opmental	Biology	
INSTITUTE AND	on Cellular Di LOCATION , Bethesda, Ma				
TOTAL MAN-YEA	2.0	PROFESSIONAL	1.7	OTHER 0.	. 3
□ (a2)	, ,	(b) Humar		(c) Neither	

This project investigates the expression of proto-oncogenes during the differentiation of embryonic lens epithelial cells to form lens fiber cells, and seeks to determine the specific function of the corresponding gene products in the developing lens. Using radioactivity labeled DNA probes we have shown that mRNA levels for two proto-oncogenes, c-myc and c-src, are specifically regulated during the differentiation of chicken embryo lens epithelial cells to form lens fiber cells, both in vivo and in vitro. Levels of c-myc mRNA are transiently elevated during the first few hours after the initiation of differentiation in vitro, as the differentiating cells withdraw from the cell cycle. Since the c-myc gene product is a nuclear, DNA-binding protein, the transient elevation of c-myc mRNA in differentiating lens cells may lead to the regulation of other, differentiation-specific genes. Levels of c-src mRNA rise more slowly than levels of c-myc mRNA, and remain elevated in the mon-dividing embryonic fiber cells. The c-src gene product is a tyrosine-specific protein kinase, the substrates of which we are presently investigating in the developing embryonic chicken lens.



PROJECT NUMBER

Z01 EY 00132-05 LMDB

PERIOD COVERE	D						
October 1, 1985 to September 30, 1986							
TITLE OF PROJEC	CT (80 characters or less	Title must fit on one	line between	the borde	rs)		
Molecular	Biology of Ph	notopigment:	S				
PRINCIPAL INVES	TIGATOR (List other prof	fessional personnel b	elow the Prin	cipal Inves	ligator) (Name title laboratory	and institute affiliation	
PI:	Toshimichi Sh	ninohara	Ph.D.		Head	LMDB, NEI	
Others:	Graeme Wistow	~	Ph.D.		Visiting Fellow	LMDB, NEI	
	Albine Katia:	l	Ph.D.		Staff Fellow	LMDB, NEI	
-	Cheryl Craft		Ph.D.		Guest Worker	LDN, NICHD	
	Masahiko Tsud	da	M.D.,	Ph.D.	Visiting Fellow	LMDB, NEI	
	Theo Van Veer	า	Ph.D.		Guest Worker	LRCMB, NEI	
COOPERATING U	NITS (d any)						
See next	page						
LAB/BRANCH							
Laborator	y of Molecular	and Develo	opmenta:	l Biol	ogy		
SECTION							
Section of	n Molecular Bi	lology					
INSTITUTE AND L	OCATION Bethesda, Mar	vland 2089	12				
TOTAL MAN-YEAR	•	PROFESSIONAL			OTHER		
TOTAL MAINTEAN	3.4	3.1	1		J. T. C.	0.0	
CHECK APPROPR						0.0	
	· · · - ·	(b) Human	ticcue		(c) Neither		
	☐ (a) Human subjects ☐ (b) Human tissues ☐ (c) Neither ☐ (a1) Minors						
	nterviews						
					d)		
SUMMARY OF WO	ORK (Use standard unred	iucea type: Do not ex	tceed the Spa	ice provide	<i>u j</i>		

We have examined the structure, function, development evolution, epitopes and uveitogenic site of the retinal S-antigen (48k protein). The cDNAs of S-antigen have been isolated from bovine retina libraries and their DNA sequences have been determined. The predicted amino acid sequence was matched completely with that of S-antigen determined by Edman degradation method. S-antigen polypeptide has some sequence homologies with Ga-transducin including the cholera toxin, pertussis toxin and enterotoxin ADP-ribosylation sites and purine nucleotide-binding sites. S-antigen is found to be a glycoprotein and its sugar moeity is glucose and mannose. Secondary structure prediction and circular dichroism analysis indicated that S-antigen has a predominantly extended (β-sheet) structure with a small C-terminal helical region. The location of the two monoclonal antibodies binding sites and the one uveitogenic site of S-antigen has been determined. These three linear immunogenic sites were localized within 20 amino acid residues at three separated sites. The mRNA of S-antigen is approximately 2 kb long and present in the rod cells but not present in the majority of cone cells. The majority of mRNA was found very close to the nucleus but not found in the most part of myoid and ellipoid of the rod inner segments. Also S-antigen mRNA was present in certain types of pinealocytes. S-antigen has only one gene in mouse and human, indicating that the S-antigen of retina and pineal is the same. The S-antigen genes of human and mouse were isolated and identified by partial DNA sequences and the human S-antigen gene is located on the chromosome No. 14.

	i.e	

SUMMARY OF WORK (Use standard unreduced type. Do not exceed the space provided.)

genetic engineering in the visual system.

PROJECT NUMBER

201 EY 00126-05 LMDB

October 1	, 1985 to Sept	ember 30, 1	986				
	CT (80 characters or less						
Crystalli	n Genes: Stru	icture, Organ	nization, E	Expression, and Evolu	ition		
	PRINCIPAL INVESTIGATOR (List other professional personnel below the Principal Investigator) (Name liftle laboratory and institute attiliation)						
PI:	Joram Piatigo	rsky	Ph.D.	Chief	LMDB, NEI		
	Ana B. Chepel	insky	Ph.D.	Expert	LMDB, NEI		
	Graeme J. Wis	tow	Ph.D.	Visiting Associate	LMDB, NEI		
-	John F. Kleme	nt	Ph.D.	Staff Fellow	LMDB, NEI		
	Mark A. Thomp	son	Ph.D.	Staff Fellow	LMDB, NEI		
	Charlotte A.		Ph.D.	Guest Worker	LMDB, NEI		
	Eric F. Wawro	usek	Ph.D.	Staff Fellow	LMDB, NEI		
See next LAB/BRANCH Laborator		and Develop	omental Bic	olgy			
SECTION	n Molecular Ge	•					
NEI, NIH,	LOCATION Bethesda, Mar	yland 20892	2				
TOTAL MAN-YEA	13.0	PROFESSIONAL	13.0	OTHER 0.0			
(a1)	RIATE BOX(ES) an subjects Minors Interviews	☐ (b) Human	tissues	(c) Neither			

We have continued to characterize the structure, expression and evolution of crystallin genes of the eye lens. Sequences have been obtained for the BB1-, BA3/A1- and B2-crystallin chicken cDNAs. Gene sequences have been derived for chicken and human aA-crystallin chicken BB1- and BA3/A1-crystallin, and chicken δ-crystallin. Both chicken δ-crystallin polypeptides were shown to be generated from the 61 mRNA by a translational or co-translational mechanism. Transfection experiments demonstrated that the alternative RNA splicing of the murine aA-crystallin gene is neither tissue- nor species-specific. Crystallin promoters were analyzed by fusion to the bacterial chloramphenical acetyl transferase (CAT) gene in the pSVO-CAT expression vector. Cell-free transcription experiments using a Hela cell extract was used to identify the core promoter of the chicken δ - and murine αA -crystallin promoters. Transient transfection experiments using cultured lens epithelia and production of transgenic mice demonstrated tissue-specific and developmental controls operating in the crystallin promoters. Both positive and putative negative regulatory sequences were indicated. The murine aA-crystallin promoter (sequences -364 to +45) when fused to the SV40 T-antigen gene neoplastically transformed lens cells in transgenic mice. Thus, these experiments initiate

DEPARTMENT OF HEALTH AND HUMAN SERVICES - PUBLIC HEALTH SERVICE

PROJECT NUMBER

	NOTICE OF INTR	AMURAL RESEARC	H PROJECT	
				Z01 EY 00078-09 LOP
PERIOD COVER	ED			
	1, 1985, to Sep			
TITLE OF PROJE	ECT (80 characters or less T	tle must lit on one line betwee	en the borders)	
		Dystrophies and		
PRINCIPAL INVE			ncipal Investigator) (Name, title, las	
PI:	Merlyn M. Rodr	igues M.D.,Ph.D	. Head, Section Ophthalmic Pat	
Others:	Joseph Hackett	B.S.	Biologist	LOP, NEI
	Reginald Gaski		Histologist	LOP, NEI
COOPERATING Departme		logy, University	of Iowa, Iowa City	
LAB/BRANCH				
Laborato	ry of Ophthalmi	c Pathology		
SECTION				
	on Ophthalmic Pa	athology		
INSTITUTE AND	LOCATION			
		NIH, Bethesda,		
TOTAL MAN-YEA	ARS P	ROFESSIONAL	OTHER	
	0.2	0.1	0.1	
CHECK APPROP (a) Hum (a1) (a2)	an subjects	(b) Human tissues	(c) Neither	

SUMMARY OF WORK (Use standard unreduced type. Do not exceed the space provided.)

Human corneal dystrophies and degenerations which have been clinically documented are studied as keratoplasty specimens with histochemical stains, scanning and transmission electron microscopy, and immunologic techniques in an attempt to elucidate pathogenetic mechanisms. This approach has provided insight into cellto-cell relationships in the normal and diseased states. In patients with primary and recurrent macular corneal dystrophy, intercellular and extracellular accumulation of fibrillogranular material was observed in the corneal stroma, Descemet's membrane, and corneal endothelium. The presence and production of collagen, glycoconjugates, and collagenase have been investigated with immunofluorescent electrophoretic, and chromatographic methods. The lectin binding patterns were compared in corneas from patients with macular dystrophy and control. The characterization of amyloid in lattice corneal dystrophy and corneal amyloid degeneration was performed using immunohistochemical stains and biochemical analysis. Lack of AA reactivity was observed in corneal amyloid deposits. Keratoplasty specimens from granular corneal dystrophy and controls were examined by combinations of immunohistological stains, transmission electron microscopy, and SDS gel electrophoresis. In granular dystrophy, the deposits consisted of phospholipid with microfibrillar protein at the edges. Corneal buttons from patients with Fuchs' dystrophy had varying degrees of clinical edema measured in most cases by preoperative optical ultrasonic pachymetry. Histologically, marked thickening of Descemet's membrane and abnormal corneal endothelium corresponded to areas of severe clinical edema and were usually located in the central and paracentral regions. Clinical edema was not present unless accompanied by marked thickening of Descemet's membrane with multiple guttata and attenuation of corneal endothelium. The peripheral cornea was relatively clear clinically and showed minimal histologic changes.

PROJECT NUMBER

ZO1 EY 00096-08 LOP

PERIOD COVERE	D						
October 1	1, 1985, to Sept	ember 3	30, 1986				
	CT (80 characters or less Tit						
Clinicopa	athologic Studie	s of Hu	man Ocular	Dise	ases		
PRINCIPAL INVES	STIGATOR (List other profess	ional personi	nel below the Princi	pal Invest	igator) (Name title laboratory and	institute a	Miliation)
PI:	Merlyn M. Rodr	igues	M.D., Ph.D	. He	ad, Section on	LOP,	NEI
	1.0. 17.1 11.	-6		Op	hthalmic Pathology		
Others:	David Bardenst	ein	M.D.	Gu	est worker	LOP,	NEI
	Joseph Hackett		B.S.	Bi	ologist	LOP,	NEI
	Reginald Gaski			Hi	stologist	LOP,	NEI
COOPERATING U	INITS (if any)						
LAB/BRANCH							
Laborato	ry of Ophthalmic	Patho:	logy				
SECTION							
Section	on Ophthalmic Pa	atholog:	y				
INSTITUTE AND L	OCATION	-					
National	Eye Institute,	NEI, B	ethesda, MI	2020)5		
TOTAL MAN-YEAR		ROFESSIONA			OTHER		
0.	3		0.2		0.1		
CHECK APPROPRIATE BOX(ES)							
☐ (a) Human subjects ☐ (b) Human tissues ☐ (c) Neither							
☐ (a1) I							
☐ (a2) I	Interviews						
SUMMARY OF WORK (Use standard unreduced type. Do not exceed the space provided.)							

Patients with localized ocular diseases or with ocular manifestations of systemic disease are examined clinically, and photographic documentation is made of significant findings. Biopsy specimens or autopsy eyes from these patients are examined by electron microscopy and histochemical stains. Studies are

performed on patients with ocular manifestations of systemic diseases.

Forty patients with acquired immunodeficiency syndrome (AIDS) were examined for ocular abnormalities. Twenty of these patients died and the eyes were obtained for culture and histologic examination. These patients have multiple opportunistic infections and neoplasms as the result of a severe depression of cellular immunity. Fifty percent of all patients with AIDS and 75% of the autopsy group have ocular signs attributable to AIDS. Ocular findings were confined to four major categories: cytomegalovirus (CMV) retinitis (10 patients), retinal cotton wool spots (11 patients), conjunctival Kaposi's sarcoma (2 patients), and neuro-ophthalmic motility abnormalities (3 patients). Cytomegalovirus retinitis was a significant cause of visual loss. Seven of 40 autopsy eyes had hand-motion or worse vision prior to the patient's death because of CMV and progressed to involve the entire retina in three to six months resulting in a gliotic retina membrane. Disseminated systemic histoplasmosis was observed in a patient with AIDS. In 3 patients, the effect of argon laser treatment was shown to be ineffective in halting the spread of cytomegalovirus in patients with AIDS.

Immunohistochemical stains are performed on patients with retinitis pigmentosa and retinoblastoma to test for the presence of neuronal and glial proteins. Electron microscopy is also performed in selected cases.



PROJECT NUMBER

ZO1 EY 00114-06 LOP

PERIOD COVER	RED			
October	1, 1985, to September	30, 1986		
TITLE OF PROJ	ECT (80 characters or less. Title must f.	t on one line between the	borders)	
Histopat	thologic Studies of Ar	nimal Models of	Human Ocular Disea	ases
PRINCIPAL INVE	ESTIGATOR (List other professional per	sonnë! below the Principal	Investigator) (Name, title, laborator	y and institute affiliation)
PI:	Merlyn M. Rodrigues	M.D.,Ph.D.	Head, Section on Ophthalmic Patholo	•
Others:	Reginald Gaskins		Histologist	LOP, NEI
•	Joseph Hackett	B.S	Biologist	LOP, NEI
	Anastasios Halkias	M.D.	Fogarty Fellow	LOP, NEI
	Barbara Wiggert	Ph.D.	Research Chemist	LVR, NEI
	Gerald Chader	Ph.D.	Chief, Laboratory of Vision Research	·
LAB/BRANCH				
Laborato	ory of Ophthalmic Path	nology		
SECTION				
	on Ophthalmic Patholo	DEY		
INSTITUTE AND	LOCATION			
	Eye Institute, NIH,			
TOTAL MAN-YE	ARS PROFESSI	ONAL	OTHER	
0,2		0.1	0.1	
	PRIATE BOX(ES)			
		luman tissues	(c) Neither	
(a1)				
□ (a2)	Interviews			

SUMMARY OF WORK (Use standard unreduced type. Do not exceed the space provided.)

Immunocytochemical staining of fresh frozen rhesus monkey retinas was performed using indirect immunofluorescence and immunoperoxidase (avidin-biotincomplex). Affinity-purified antibodies to interphotoreceptor retinoid-binding protein (IRBP) obtained from rabbits was used to localize IRBP on frozen sections. Fresh frozen pineal glands from the same species were stained by the avidin-biotin-peroxidase method. In addition, retinas from rod-dominant and cone-dominant species were examined. Immunocytochemical staining revealed localization of IRBP in the interphotoreceptor space of peripheral equatorial and posterior retina, with marked decrease in staining in the fovea. A transition zone was noted at the ora serrata, where staining was present in the peripheral retina up to the ora serrata, but was absent in ciliary epithelium. Cone-dominant retinas (chick and turtle) showed lack of reactivity to IRBP. Rod-dominant rat retina showed localization of IRBP to the interphotoreceptor space. Primate and rat pineal showed immunocytochemical localization of IRBP. Spontaneously occurring anterior chamber segment anomalies in DBA/2 mice were studied by slit-lamp biomicroscopy and light and transmission electron microscopy (TEM). The opacities consisted of aggregates of basophilic material in the superficial stroma which stained positively for elastin TEM revealed that they were electron dense and extracellular. Iris abnormalities consisted of stromal atrophy and proliferation of corneal endothelium and basement membrane across the iris surface and trabecular meshwork. The corneal opacities seen in DBA/2 mice show a striking similarity to those which characterize familial band~ shaped nodular keratopathy, a form of corneal elastosis.

ANNUAL REPORT NATIONAL EYE INSTITUTE October 1, 1985 - September 30, 1986

REPORT OF THE CHIEF, LABORATORY OF RETINAL CELL AND MOLECULAR BIOLOGY Gerald J. Chader, Ph.D.

The Laboratory of Retinal Cell and Molecular Biology is a newly formed unit within the National Eye Institute. It draws together former members of the Laboratory of Vision Research who have a common interest in basic and applied studies on the neural retina and associated tissues such as pigment epithelium. As the name implies, aspects of retinal cell biology as well as molecular biology are being studied both in normal and in diseased tissues. In particular, there is an emphasis on the study of inherited retinal diseases as exemplified by retinitis pigmentosa and retinoblastoma.

The laboratory consists of three separate but cooperating Sections each investigating a particular aspect of retinal function. Following are highlights of some of the research conducted by members of the LRCMB over the last year:

Section on Cell Biology: As Chief of this section, Dr. Paul O'Brien leads the NEI effort in studying unique biochemical events in the retina related to photoreceptor physiology. In normal retinas, he has continued to define specific reactions leading to acylation of rhodopsin, the visual pigment. Palmitate is added to newly synthesized as well as mature rhodopsin molecules and may be involved in important cellular processes such as sorting and transport of protein from the Golgi apparatus and in disc membrane assembly and outer segment membrane repair mechanisms. In parallel, he has been investigating synthesis and degradation of proteins and phospholipids in normal retinas and in animal models of inherited retinal degeneration. Of potential importance is the apparently abnormal phospholipid metabolism that Dr. O'Brien has now observed in poodles affected with inherited rod-cone degeneration. These animals exhibit an abnormally low rate of photoreceptor outer segment renewal prior to the onset of degeneration.

Section on Biochemistry: Dr. Barbara Wiggert has continued her impressive work on the biochemistry and function of IRBP, the interphotoreceptor retinoid-binding protein. It is now clear that this protein is a unique, new type of retinoid transport protein designed for the extracellular transport of retinoids between the photoreceptor elements of the neural retina and vitamin A stores in the retinal pigment epithelium. This work takes on added significance with the discovery of substantial amounts of IRBP in the pineal organs of several species. Although the function of the protein in the pineal gland in presently not clear, it may indicate a hitherto unknown role of retinoids and their transport in this secondary organ of phototransduction.

Section on Gene Regulation: In collaboration with Dr. Wiggert, Dr. John Nickerson and his group have made excellent progress in studying the molecular biology of the IRBP molecule. cDNAs for IRBP, for example, have now been established and verified and about one half of the nucleotide sequence has been determined. Bovine and human genomic clones have also been established and progress has been made concerning the chromosomal location of the IRBP gene. Aspects of IRBP synthesis are also under investigation. It is of interest that the size of the IRBP mRNA is extraordinarily large, indicating a long uncoded region. In situ hybridization studies have indicated that the probable site of IRBP synthesis in the retina is predominantly in the rod photoreceptor cell. These studies further strengthen the argument that IRBP is intimately involved in general processes of photoreception.

Other members of the Section on Gene Regulation have focused their work on attempting to find the biochemical lesion(s) in animals exhibiting early-onset retinal degeneration. In particular, Mr. R.T. Fletcher has studied genetic crosses of rd and rds mice with reference to the cyclic GMP second messenger system in photoreceptor cells and has found a direct gene-dose relationship between a photoreceptor cell cyclic GMP phosphodiesterase enzyme and the rds gene. This would seem to be of importance in linking abnormal retinal concentrations of cyclic GMP with the rds gene responsible for degeneration of the retina.

Another hereditary disease under investigation is retinoblastoma. Dr. A. Kyritsis has made substantial progress in understanding factors related to growth and differentiation of human Y-79 retinoblastoma cells in culture. One of the most controversial questions concerning this type of cancer over the years, has been the origin of the cells. From Dr. Kyritsis' work, it now appears that the cells are multipotential blast cells capable of at least partial differentiation into most if not all of the general cell types found in the normal retina. These include neuronal, glial and pigment epithelial elements. This has been a particularly satisfying finding, since it explains most of the seemingly contradictory previous evidence in the literature and it indicates that the tumor cells, depending on their particular surroundings (substratum, differentiating agents, etc.), can develop along any one or more of these pathways as the tumor grows.

PROJECT NUMBER

Z01 EY 00070-09 LRCME

PERIOD COVE	RED					
October	1, 1985 to Septem	per 30, 1986				
PI: Barbara Wiggert Ph.D. Head, Section on Biochemistry Others: Ling Lee M.S. Chemist Michael Redmond Ph.D. Staff Fellow Gerald J. Chader Ph.D. Chief COOPERATING UNITS (# any) LSU Eye Center, New Orleans, LA (N. Bazan, T. Reddy)						
October 1, 1985 to September 30, 1986 TITLE OF PROJECT (80 characters or less Title must fit on one line between the borders) Vitamin A and Ocular Tissues PRINCIPAL INVESTIGATOR (List other professional personnel below the Principal Investigator) (Name title laboratory and institute at PI: Barbara Wiggert Ph.D. Head, Section on LR Biochemistry Others: Ling Lee M.S. Chemist LR Michael Redmond Ph.D. Staff Fellow LR Gerald J. Chader Ph.D. Chief LR COOPERATING UNITS (If any) LSU Eye Center, New Orleans, LA (N. Bazan, T. Reddy) LAB/BRANCH Laboratory of Retinal Cell and Molecular Biology						
PRINCIPAL IN	VESTIGATOR (List other profess	iona personnel below the Prin	cipa Investigator) (Name	title laboratory and	institute affiliation	
PI:	Barbara Wiggert	Ph.D.	·		LRCMB,	NEI
Others:	Ling Lee	M.S.	Chemist		LROMB,	NEI
	Michael Redmond	Ph.D.	Staff Fell	. OW	LRCMB,	NEI
	Gerald J. Chades	Ph.D.	Chief		LROMB,	NEI
	, , ,	ns, LA (N. Bazan	, T. Reddy)			
LAB/BRANCH						
Laborato: SECTION	ry of Retinal Cel	l and Molecular 1	Biology			
Section (on Biochemistry DLOCATION					
NEI, NIH	Bethesda, Maryla	and 20892				
TOTAL MAN-Y	EARS PF	ROFESSIONAL	OTHER			
	2.7	1.7		1.0		
	OPRIATE BOX(ES)					
	man subjects 🗵	(b) Human tissues	☐ (c) Neith	ner		
) Minors					
☐ (a2) Interviews					
SUMMARY OF	WORK (Use standaro unreduce	d type. Do not exceed the spi	ace provided)			

An enzyme-linked immunosorbent assay (ELISA) for monkey and human Interphoto-receptor Retinoid-Binding Protein (IRBP) was used to quantitate IRBP in normal and diseased human retinas and retinoblastoma tumors. IRBP levels were uniformly low in retinas from human cases of hereditary retinal degenerations even in areas in which photoreceptors remained. IRBP was present in several retinoblastoma tumors examined.

The amino terminal sequences of monkey and bovine IRBPs were extended to over 30 residues each. The major monkey sequence had an additional 5 amino acid residues at its amino terminus not observed with bovine IRBP, although the sequences showed extensive homology. The amino terminal sequence of human IRBP was identical to that of the monkey, and the two sequences were present in equal amount.

PROJECT NUMBER

Z01 EY 00015-21 LRCMB

					1		
					lers)		
PERIOD COVERED October 1, 1985 to September 30, 1986 TITLE OF PROJECT (80 characters or less Title must fit on one line between the borders) The Cell Biology of the Vertebrate Retina PRINCIPAL INVESTIGATOR (List other professional personnel below the Principal Investigator) (Name title laboratory and institute affiliation) PI: Paul J. O'Brien Ph.D. Head, Section on LRCMB, NEI Cell Biology Others: Robert St. Jules Ph.D. Staff Fellow LRCMB, NEI COOPERATING UNITS (M any) Department of Anatomy, University of Toronto (M. J. Irons) LABIBRANCH Laboratory of Retinal Cell and Molecular Biology Section on Cell Biology INSTITUTE AND LOCATION NEI, NIH, Bethesda, Maryland 20892							
PRINCIPAL INVES	TIGATOR (List other prof	'essional perso	nnel below the P	rıncıpa! Inve	stigator) (Name title laboral	tory and institute affil	liation)
PI:	Paul J. O'Bri	en l	Ph.D.	Head,	Section on	LRCMB,	NEI
					Cell Biology		
-							
Others:	Robert St. Ju	iles I	Ph.D.	Staff	Fellow	LRCMB,	NEI
COOPERATING U	NITS (d any)						
Department	of Anatomy,	Uni versi	ity of Tor	ronto	(M. J. Irons)		
	of Retinal C	ell and	Molecular	Biolo	у		
NEI, NIH,	Bethesda, Mar	yland 20	892				
TOTAL MAN-YEAR	15	PROFESSION	VAL.		OTHER		
	1.6		1.6		0	.0	_
CHECK APPROPR							
(a) Huma	in subjects	□ (b) Hu	ıman tissues	s 🖸	(c) Neither		
☐ (a1) N	Minors						
☐ (a2) I	nterviews						
CHILDY OF MIC	DV 41				la of 1		

SUMMARY OF WORK (Use standard unreduced type. Do not exceed the space provided.)

The acylation of rhodopsin with palmitic acid has been studied to determine the function of this modification in the intracellular transport of rhodopsin and its incorporation into functional disc membranes. Palmitate is added to rhodopsin shortly after the polypeptide is synthesized. Most of the new rhodopsin molecules are retained in the endoplasmic reticulum for at least six hours, suggesting that transport to the outer segment may not be a continuous process. Palmitate is also added to mature rhodopsin molecules in the outer segment indicating a possible repair mechanism or a physiologically important cycle of removal and replacement. The phospholipids of the outer segment turn over continuously but retain and, in the case of phosphatidylethanolamine, even increase the level of labeling with palmitic acid. Thus, this fatty acid undergoes a continuous process of reutilization.

Cytidine monophosphate, a by-product of phospholipid turnover, is cleaved by a manganese-dependent nucleotidase in rod outer segments. Histochemical studies indicate that this enzyme is transported from the pigment epithelium to the outer segments in anticipation of disc shedding and may represent a class of degradative enzymes involved in this process.



PROJECT NUMBER

IO1 EY 00016-19 LRCMB

PERIOD COVERE	D						
October 1	, 1985 to Sept	ember 30,	1986				
TITLE OF PROJEC	T (80 characters or less	Title must fit on o	ne line betwee	n the bord	ders)		
The Bioche	emistry of Nor	mal and Dy	strophi	c Ret	inas		
PRINCIPAL INVES	TIGATOR (List other pro-	essional personnel	below the Pri	ncipal Inve	stigator) (Name title labora:	tory and institute af	filetion
PI:	Paul J. O'Bri	en Ph.	D. :	Head,	Section on Cell Biology	LRCMB,	NEI
Others:	Peter A. Dudl	ey Ph.	D.			ECP,	NEI
		dicine, Ur	niversit	y of I	Pennsylvania (G.	Aguirre)	
LAB/BRANCH							
	of Retinal C	ell and Mo	lecular	Biol	DRY		
SECTION							
	n Cell Biology						
INSTITUTE AND L	OCATION						
	Bethesda, Mar						
TOTAL MAN-YEAR	RS	PROFESSIONAL			OTHER		
	0.2		0.2_		0	.0	
CHECK APPROPR							
🗌 (a) Huma	•	🗀 (b) Huma	an tissues	L	🗓 (c) Neither		
_ ' '	Minors						
□ (a2) I	nterviews						
SUMMARY OF WO	DRK (Use standard unred	ucad type. Do not	exceed the sp	ace provid	ted)		

This project examines biochemical events unique to the retina, particularly the synthesis and modification of photoreceptor membrane components, in the retinas of vertebrates which can be affected by inherited retinal degenerations. The synthesis of the visual pigment, rhodopsin, occurs at a normal rate as measured by radioactive leucine incorporation following intravitreal injection in the eyes of miniature poodles affected with progressive rod-cone degeneration. Similarly, the glycosylation and acylation of rhodopsin were found to be normal following intravitreal injection of labeled fucose or palmitic acid, respectively. However, phospholipid synthesis or degradation, measured by radioactive palmitic acid incorporation, appears to be different in the affected dogs, suggesting a possible metabolic defect in this inherited disorder.

PROJECT NUMBER

Z01 EY 00148-13 LRCMB

PERIOD COVER	RED					
October 1	, 1985 to Septe	mber 30, 19	986			
PERIOD COVERED October 1, 1985 to September 30, 1986 TITLE OF PROJECT (80 characters or less Title must fit on one line between the borders) Cyclic Nucleotides and Visual Control Mechanisms PRINCIPAL INVESTIGATOR (List other professional personnel below the Principal Investigator) (Name little laborator, and institute affiliation). PI: Gerald J. Chader Ph.D. Chief LRCMB, NEI Others: Susan Gentleman Ph.D. Expert LRCMB, NEI R. Theodore Fletcher M.S. Chemist LRCMB, NEI Robert L. Somers B.S. Chemist LRCMB, NEI C. Lal Kapoor Ph.D. Guest Worker LRCMB, NEI COOPERATING UNITS (if any) Section on Medical Genetics, School of Veterinary Medicine, University of Pennsylvania (G. Aguirre); Department of Anatomy, Erasmus University, Rotterdam, The Netherlands (S. Sanyal) LABBERANCH Laboratory of Retinal Cell and Molecular Biology SECTION						
Cyclic Nu	cleotides and V	isual Contr	rol Mechani	sms		
					laboraton, and institute a	affiliation)
PI:	Gerald J. Chad	er	Ph.D.	Chief	LRCMB,	NEI
Others:	Susan Gentlema	n	Ph.D.	Expert	LRCMB,	NEI
	R. Theodore Fl	etcher	M.S.	Chemist	LROMB,	NEI
•	Robert L. Some	rs	B.S.	Chemist	LRCMB,	NEI
	C. Lal Kapoor		Ph.D.	Guest Worker	LRCMB.	NEI
COOPERATING	LINITS (d apv)					
	* *					
			ent of Anat	omy, Erasmus l	Jniversity, Ro	otterdam,
	rlands (S. Sany	al)				
	y of Retinal Ce	ll and Mole	ecular Biol	ogy		
	n Gene Regulati	on				
INSTITUTE AND						
NEI, NIH,	Bethesda, Mary	land 20892	2			
TOTAL MAN-YE	ARS	PROFESSIONAL		OTHER		
	2.8		1.8		1.0	
	PRIATE BOX(ES)					
	nan subjects	$\overline{\mathbb{X}}$ (b) Humar	i tissues	(c) Neither		
	Minors					
(a2)	Interviews					
SUMMARY OF V	NORK (Use standard unredu	iced type. Do not ex	ceed the space pr	ovided)		

Cyclic nucleotides and protein phosphorylation play a special role in photo-receptor control mechanisms.

- 1. A calcium phospholipid-dependent protein kinase (C-kinase) phosphorylates specific proteins of the rod outer segment organelle.
- 2. A specific extracellular cyclic CMP phosphodiesterase was identified in the retinal extracellular matrix.
- 3. Cyclic AMP-dependent protein kinases were found to be abnormal in human retinoblastoma cells in culture.
- 4. Cyclic GMP was found to be abnormally high in photoreceptor cell synapses of an animal model of inherited retinal degeneration.

PROJECT NUMBER

Z01 EY 00124-06 LRCMB

PERIOD COVER	ED					
October 1	, 1985 to Sept	tember 30, 1	986			
TITLE OF PROJ	ECT (80 characters or less	Title must fit on one	e line between th	ne borders)		
Metabolis	m of the Retir	a and Pigme	nt Epithe	elium		
PRINCIPAL INVE	STIGATOR (List other pro	ofessional personnel t	elow the Princip	a Investigator) (Name title labo	oraton, and institute a	Mulation)
PI:	Gerald J. Cha	ader	Ph.D.	Chief	LRCMB,	NEI
Others:	Shay-Whey M.	Koh	Ph.D.	Staff Fellow	LRCMB,	NEI
	Athanassios F	. Kyritsis	M.D.	Visiting Fellow	LRCMB,	NEI
-	R. Theodore F	letcher	M.S.	Chemist	LRCMB,	NEI
COOPERATING	UNITS (if any)					
LAB/BRANCH						
Laborator	y of Retinal C	Cell and Mol	ecular Bi	lology		
SECTION						
Section o	n Gene Regulat	ion				
INSTITUTE AND	LOCATION					
NEI, NIH,	Bethesda, Mar	yland 2089	2			
TOTAL MAN-YE	ARS	PROFESSIONAL		OTHER		
	2.1		2.1		0.0	
CHECK APPROP						
		🔀 (b) Humar	n tissues	(c) Neither		
☐ (a1)						
	Interviews					
SUMMARY OF V	VORK (Use standard unred	duced type. Do not e.	xceed the space	provided)		

Metabolic characteristics of retinal cells in tissue culture are under examination.

- 1. The response of cultured cells of the retinal pigment epithelium to various neurotransmitters and neuromodulators have been examined, especially to vasoactive intestinal peptide (VIP). A specific VIP receptor has been identified on pigment epithelial cell membranes; also, soluble and membrane proteins phosphorylated in response to VIP have been identified.
- 2. Human retinoblastoma cells in culture can be induced to differentiate in culture into the three major cell types seen in the normal retina, ie, neurons, glia and pigment epithelial cells. Laminin and appropriate other attachment factors appear to play a role in retinoblastoma cell differentiation as well as attachment.

*		

DEPARTMENT OF HEALTH AND HUMAN SERVICES - PUBLIC HEALTH SERVICE

PROJECT NUMBER

	NOTICE OF INT	RAMURAL RES	EARCH PROJE	:CT	Z01 E	Y 00196-03	3 LRCMB
PERIOD COVER	ED						
October 1.	, 1985 to Septe	ember 30, 198	6				
TITLE OF PROJE	CT (80 characters or less	Title must fit on one hi	e between the borde	5)			
Molecular	Genetics of the	ne Eye and Oc	ular Diseas	es :			
PRINCIPAL INVE	STIGATOR (List other pro	lessional personnel belo	w the Principal Inves	igator) (Name	title laboratory and a	nstitute at lation)	
PI:	John M. Nicker	rson P	h.D. Se	nior Staf	f Fellow	LRCM5,	NEI
Others:	Diane Borst	P	h.D. Gu	est Worke	r	LRCMB.	NEI
	Shirley Rainie	er P	h.D. St	ff Fello		LRCMB.	
-	T. Michael Red					LRCMB,	
COOPERATING DOORS DO	UNITS (d any) epartment, Univ	versity of Lu	nd, Lund, S	weden (Th	eo Van Veen)	
LAB/BRANCH							
Laboratory	of Retinal Co	ell and Molec	ular Biolog	,			
SECTION							
Section or	n Gene Regulati	on					
INSTITUTE AND	LOCATION						
NEI, NIH,	Bethesda, Mary	land 20892					
TOTAL MAN-YEA	RS	PROFESSIONAL		OTHER			
	2.5		2.5		0.0		
CHECK APPROP							
	an subjects		issues \square	(c) Neithe	r		
(a1)							
□ (a2)	Interviews						

SUMMARY OF WORK (Use standard unreduced type. Do not exceed the space provided.)

My laboratory is presently isolating and characterizing recombinant DNA molecules necessary for the study of the structure and expression of IRBP (Interphotoreceptor Retinoid-Binding Protein). Clones and partial sequences from bovine IRBP have been obtained. We have cloned and completely sequenced six overlapping cDNA clones that constitute about 3500 bases of the 7000 base long bovine IRBP mRNA. The IRBP mRNA is unusually long especially given that the polypeptide encoding region of the mRNA should only require about 3600 bases. There is only one band on a Northern blot suggesting that there is only one IRBP mRNA species. The clones were used as probes for in situ hybridizations to frozen sections of bovine and monkey retinas. In large, only the rod cell perikarya were labelled with the probe. This showed us that the gene is regulated and expressed only in the rod cell, and it defined the site of synthesis of the IRBP polypeptide as the rod cell. Part of the gene for bovine IRBP has been cloned. Partial DNA sequence analysis revealed a perfect match of 75 bases of the cDNA and genomic clones. The sequences diverged at a consensus splice acceptor site in the gene clone, indicating an intron-exon boundary. We have obtained a putative genomic clone for the human gene, but it remains to be proven (by DNA sequencing) that this clone is authentic. We have begun to characterize the IRBP gene by analysis of Southern genomic blots. We have shown that the gene is present in only one copy per haploid genome in a variety of species. In human, the gene is not on the X-chromosome, since hybridizing bands from male and female DNA samples are the same intensity. We are now in the process of mapping the precise chromosome.



ANNUAL REPORT NATIONAL EYE INSTITUTE October 1, 1985 - September 30, 1986

REPORT OF THE CHIEF, LABORATORY OF SENSORIMOTOR RESEARCH Robert H. Wurtz, Ph.D.

This is the eighth Annual Report of the Laboratory of Sensorimotor Research and is intended to describe work in the Laboratory over the last two years. I would like in this somewhat more flexible report to concentrate on the relevance of work in the Laboratory to several fundamental problems of visual neuroscience. In particular, I would like to emphasize the contribution of our analysis of the visual and the oculomotor system in the monkey to an understanding of these systems in man.

The aspect of brain function that we study, the visual and oculomotor system, has been shown repeatedly to be very similar in man and monkey so that the experiments we perform on the monkey serve as a model for man. Behavioral, physiological, and anatomical experiments possible on the monkey have given us our most fundamental understanding of how the visual and oculomotor functions of the brain of man are likely to be organized. In addition I think several investigations in the Laboratory in the last several years illustrate how the precise analysis possible in the visual-oculomotor system has allowed exploration of fundamental questions in brain research.

One of these questions is how we are able to obtain sensory information in spite of the continual movements we make in our environment. The pinnacle of this stability is the ease with which we maintain stable vision in spite of our own movement of body, head, and eyes. Much of this stability of the eye is due to the vestibular-ocular reflex, a system that Dr. F. A. Miles in the Laboratory has studied over a period of years. In this system, the balance mechanisms in the ear lead to eye movement that compensates for head and body movement. In addition to this vestibular mechanism, another system has been studied by Dr. Miles that provides fine tuning of this stabilization. This ocular following response occurs with the incredibly short latency of only about 50 msec following a rapid or saccadic eye movement across a visually patterned environment. Furthermore, the ocular following is enhanced following such eye movements just at the time when increased stabilization of vision is essential. The mechanisms underlying this visual motor event can be dissected apart by replacing the rapid eye movement with an equally rapid shift in the visual environment: the ocular following response can still be invoked easily. However, when a large visual scene is quickly moved in a saccade-like way the ocular following response is actually suppressed. This powerful inhibitory mechanism that emanates from the peripheral visual field probably prevents the eye from tracking the visual disturbances created by the rapid eye movements as they sweep the visual scene across the Thus, not one visual mechanism is involved in stabilization, but rather two interacting ones: one to produce the stabilization of the eye, the other to produce a suppression of that stabilization when it is inappropriate. experiments demonstrate graphically the series of exact and elaborate mechanisms that primates use to maintain the essential conditions for normal vision. The

90	

deficits in this type of stabilization, which may occur with the diseases of man, can only begin to be diagnosed if the series of interlocking mechanisms is clearly understood and this understanding can only be derived from the type of carefully controlled experiments such as those of Dr. Miles.

Effective vision not only demands stability during periods of fixation on a given object, but it also demands the ability to shift the highly developed central or foveal region of the retina from one part of the visual field to another. This selective function, frequently referred to as selective visual attention, has been studied over a number of years in the Laboratory. Dr. D. L. Robinson and his collaborators have continued this work in both monkey and man. vious physiological experiments had shown that a region of cerebral cortex, the posterior parietal cortex, was critically involved in tasks related to attention, and that the pulvinar nuclei of the thalamus, a step on the visual pathway reaching from the brainstem to parietal cortex, were also critically involved in this selective attention mechanism. This behavioral attention in monkeys can be studied by giving the monkey a cue indicating the area of the visual field that is important and then testing the speed of a monkey's subsequent response to a stimulus in that field. By suppressing the activity of one area area of the brain with minute injections of chemicals while the monkey performs this task, Dr. Robinson has been able to show the involvement of that area of the brain in this type of selective attention. These methods of behavior testing have now been extended to human patients whose disease might involve the areas of the brain identified in the monkey as being related to visual attention. The tests have provided a constellation of deficits that characterize patients with lesions of the parietal area of cerebral cortex. In contrast, patients suffering from schizophrenia or from Alzheimer's disease might show some deficits in attentional tasks, but not the same set of deficits seen in those patients with parietal lesions. Thus, a set of experiments growing out of an analysis of the neural systems within the brain of monkeys has been developed that allows the diagnosis of damage to one part of the brain of man. Furthermore, this analysis of visual attention shows that the highest level of behavior in man can be subjected to precise behavioral analysis and related to precise neural structures within the brain.

Possibly the region of the cerebral cortex involved in the highest neural processing is the frontal lobe, and it is in this region that Dr. M. E. Goldberg and his collaborators have concentrated their analysis of one type of eye movement, the rapid, or saccadic eye movements which I have referred to above. pathways for this particular type of movement, that connects the visual signals arriving from the eye to the activation of the eye muscle to move the eye towards the target, is probably the best understood sensorimotor pathway within the brain. What is striking about this system is that there seems to be a separate segment related to the same movement made under different conditions: some areas of the brain participate in the generation of all saccades; others participate only in saccades that are important to an animal's behavior, whether or not the saccades are made in response to visual targets. The frontal cortex seems to be particularly important for this latter type of movement. The frontal eye fields do not, however, send a motor message for the generation of saccades irrelevant to a subject's behavior. Ordinarily, the act of attentive fixation filters out the signals for irrelevant saccades. Thus Dr. Goldberg and his group found that it was more difficult to evoke saccades electrically from the frontal eye fields when a monkey was actively looking at a spot of light, and the saccade evoked had a characteristic waveform called a square-wave jerk. In collaboration with the

Neuro-Ophthalmology Section, Dr. Goldberg noticed that certain patients complaining of slow reading made square-wave jerks while they attempted to fixate a target. Since these eye movements resemble those evoked in the monkey during attentive fixation, they might in man be a sign of attentional fluctuations, which would make the oculomotor system more susceptible to signals for irrelevant saccades. Since other patients with cerebellar disease have square-wave jerks but no reading difficulty, Dr. Goldberg hypothesized that the deficit was not oculomotor instability but rather an attentional deficit which interfered both with adequate oculomotor performance and reading. A small number of these patients were treated with methylphenidate, a drug approved for attentional deficit disorders. Not only did the square-wave jerks disappear, but several measures of reading performance also improved under treatment. This promising new line of clinical research in man would not have been possible without the fundamental observations on the frontal eye fields of monkeys.

Visual processing in both man and monkey is represented in other regions of the cerebral cortex, primarily in the occipital and temporal lobes. most promising developments in our understanding of this visual processing has been the realization that the analysis might be divided between different subregions of this region of cerebral cortex. My collaborators and I have studied one aspect of this visual processing, that related to visual motion processing and its relationship to a type of eye movement that is dependent on such motion, the generation of smooth pursuit eye movements. These movements are used to follow a slowly moving target in order to keep the target positioned upon the foves. It is interesting to note that this type of eye movement evolves most clearly in primates so that in order to study the system in man, one must analyze the mechanisms in another primate such as the monkey. Our experiments have demonstrated that minute chemical removal of cells in the motion area of cerebral cortex reduces a monkey's ability to generate pursuit eye movements. Furthermore, similar damage to the next step of visual processing produces an additional deficit, a difficulty in keeping the eye on the target once it is on the fovea but only when the target moves toward the damaged side of the brain. It is striking that one of the first deficits related to eye movements in man that was specifically related to the cerebral cortex was this directed pursuit deficit. with minute lesions in monkeys, we have been able to replicate the deficits seen following the uncontrolled and extensive accidents of nature in man. Furthermore, because we are able to analyze the response of single cells in the monkey, we have been able to see the types of cells in a sequence of visual areas, including those areas producing the directional deficit. The hypothesis that we have developed suggests that the first visual area is related to visual motion processing while the next one (related to the directional deficit) has signals related to maintaining eye movements that do not depend on the visual input. These experiments demonstrate the use of a particular type of eye movement in the analysis of visual processing; the removal of a tiny visual area leads to an exactly measurable deficit in the eye movement. Tight coupling between the visual and oculomotor system makes such an analysis possible in the visual system, and holds promise of precise localization of visual function even within the cerebral cortex.

Finally, in the studies considered so far, the response of neurons has generally been taken as the total number of electrical discharges produced in response to a given visual stimulus, or in relation to a particular eye movement. However, the work of Dr. L. M. Optican in this Laboratory and Dr. B. J. Richmond in the Institute of Mental Health have called this assumption into question at



least as far as the visual analysis related to form perception is concerned. They have analyzed the distribution of electrical discharges of cerebral cortex cells following the presentation of a systematically varied visual stimulus. They find that while the rate of discharge conveys information about the visual stimulation, the amount of information conveyed by the pattern of discharges is twice that carried if one looks at the number of discharges alone. They show that in a given cell several independent patterns of spike activity are produced by the set of stimuli they used and that these patterns exist simultaneously. These experiments in turn led Optican and Richmond to formulate a new hypothesis about how information is transmitted in the visual system. They propose that these neurons each act as simultaneously active but independent spatial to temporal filters. Each filter generates a component of the visual response, and these components are then combined to form the spike train. While the underlying principle of multiple, simultaneous filtering operations has been demonstrated for neurons in the primary striate cortex and the inferior temporal cortex, the same mechanism may be functioning in other systems as well. Thus, in the case of the visual system, where the stimulation can be very exactly controlled using a mathematically definable set of stimuli, it has been possible to relate the temporal pattern of cell discharge to the sensory stimulation. The extent and importance of this type of information transmission in the visual system remains almost totally unexplored, but represents an exciting possible neuromechanism that may help us to understand the way in which the complexities of vision are represented within the brain.

Market Parket		

DEPARTMENT OF HEALTH AND HUMAN SERVICES - PUBLIC HEALTH SERVICE NOTICE OF INTRAMURAL RESEARCH PROJECT

PROJECT NUMBER

201 EY 00049-08 LSR

PERIOD COVE October	RED 1, 1985, to Septem	mber 30, 1986		
		must fit on one line between the bo	rders) 8 and Visual Attention	
			vestigator) (Nama, title, laboratory and in	
PI:	Michael E. Goldbe	erg M.D.	Chief, NMS	LSR, NEI
Others;	Mark Segraves	Ph.D.	Staff Fellow	LSR, NEI
-	Deng Shu-yi	M.D., Ph.D.	Visiting Fellow	LSR, NEI
	Rolf Boch	Ph.D.	Visiting Fellow	LSR, NEI
LAB/BRANCH Laborato	ry of Sensorimoto	Research		
SECTION Neuro-Op	hthalmologic Mecha	anisms Section		
NEI, NIH	D LOCATION , Bethesda, Maryla	and 20892		
TOTAL MAN-Y		OFESSIONAL	OTHER	
	4.9	3.8	1.1	
⊠ (a) Hu □ (a1	man subjects) Minors) Interviews	(b) Human tissues	(c) Neither	
•		I type. Do not exceed the space pro-		

By antidromic stimulation in the superior colliculus of neurons in the arcuste frontal eye fields, it was determined that cells bearing eye movement or fixation signals but not those with peripheral visual information project to the superior colliculus from the frontal cortex. This implies that the cerebral cortex sends only a motor message to the brainstem saccadic eye movement system.

Monkeys trained on a saccadic adaptation paradigm learn quickly to change the amplitude of their saccades in response to intrasaccadic stimulus steps. Stimulation of the superior colliculus in the adapted case yields the same saccades as the unadapted case. The activity of single neurons also shows no evidence of adaptation: the visual and movement signals emanating from the colliculus are the veridical signals for stimulus and movement, rather than movement signals in a visual frame which are modified by the brainstem. Thus the change in saccade amplitude which occurs as a result of adaptation is compensated for at early sites in the neural chain from stimulus to response.

A small percentage of humans with reading disorders have fixational instability manifest by saccadic intrusions or square-wave jerks while they attempt to fixate a spot of light. A single dose of methylphenidate decreases or eliminates the saccadic intrusions and results in temporary improves in both the ocular mechanics of oral reading and the reading performance itself.

DEPARTMENT OF HEALTH AND HUMAN SERVICES - PUBLIC HEALTH SERVICE NOTICE OF INTRAMURAL RESEARCH PROJECT

PROJECT NUMBER

Z01 EY 00153-04 LSR

				00133 04 L3K
PERIOD COVER October	RED 1, 1985, to Sept	ember 30, 1986		
TITLE OF PROJ	JECT (80 characters or less Ti	the must fit on one line between the rimate Oculomotor		
PRINCIPAL INV	ESTIGATOR (List other profes	sional personnel below the Princip	al Investigator) (Name title laboratory and institut	le affiliation)
PI:	Frederick A. Mi	les D.Phil	Chief, OCS	LSR, NEI
Others:	Lance Optican Reuben Gellman	Ph.D. Ph.D.	Senior Staff Fellow Visiting Fellow	LSR, NEI LSR, NEI
COOPERATING	UNITS (if any)			
Laborato	ry of Sensorimot	or Research		
SECTION Oculomot	or Control Secti	on		
NEI, NIH	, Bethesda, Mary	land 20892		
TOTAL MAN-YE	3.5	ROFESSIONAL 1.5	OTHER 2.0	
	PRIATE BOX(ES) man subjects Minors Interviews	(b) Human tissues	(c) Neither	

SUMMARY OF WORK (Use standard unreduced type. Do not exceed the space provided.)

Experiments were concerned with the initial ocular following responses to transient ramp movements of the visual scene in monkeys. These tracking movements are important in the stabilization of gaze which is so necessary for good visual acuity. We had previously shown in monkeys that responses have short latency (50 msec) and are transiently enhanced after saccadic eye movements due to the associated visual disturbance. We now report that saccadic eye movements are also associated with powerful inhibitory effects on ocular following. This inhibition was shown to be visual in origin and to result from stimulation of the peripheral region of the visual field. For these experiments the visual field was partitioned into central and peripheral regions (center, 20° diameter). Sudden shifts of the peripheral images elicited brief, powerful inhibition of ocular following responses generated by test stimuli at the center. This peripheral suppression showed good intraocular transfer: saccade-like movements of the peripheral field of one eye suppressed the responses elicited by test ramps applied at the center of the other eye. These data indicate that the suppression involves lateral spatial interactions at/or beyond a site that receives input from the two eyes, and hence, must be mediated by the central nervous system. We suggest that this suppression functions to prevent the ocular following system from tracking the visual disturbances caused by saccades: saccade-like movements of the central field alone produced small transient ocular following responses whereas such movements of the periphery or of the whole field did not.

DEPARTMENT OF HEALTH AND HUMAN SERVICES - PUBLIC HEALTH SERVICE NOTICE OF INTRAMURAL RESEARCH PROJECT

PROJECT NUMBER

201 EY 00152-04 LSR PERIOD COVERED October 1, 1985, to September 30, 1986 TITLE OF PROJECT (80 characters or less. Title must fit on one line between the borders.) Visually Induced Adaptive Changes in Saccadic Innervation PRINCIPAL INVESTIGATOR (List other professional personnel below the Principal Investigator) (Name, title, leboratory, and institute administrant PI: Lance Optican Ph.D. Res. Biomedical Engineer LSR, NEI Others: Frederick A. Miles D.Phil. Chief, OCS LSR, NEI COOPERATING UNITS (if any) David S. Zee, M.D., Professor Neurology, Johns Hopkins School of Medicine, Baltimore, MD LAB/BRANCH Laboratory of Sensorimotor Research SECTION Oculomotor Control Section INSTITUTE AND LOCATION NEI, NIH, Bethesda, Maryland 20892 TOTAL MAN-YEARS PROFESSIONAL OTHER 0.8 0.8 0.0

(a2) Interviews

SUMMARY OF WORK (Use standard unreduced type. Do not exceed the space provided.)

(b) Human tissues

CHECK APPROPRIATE BOX(ES)

(a) Human subjects

(a1) Minors

Saccades are the rapid eye movements used to change visual fixation. These eye movements are very fast yet they end abruptly, i.e., the eyes do not drift at the end of a saccade. This lack of post-saccadic drift is essential for good visual acuity after a saccade. Previous experiments in this laboratory have shown that the brain actively suppresses post-saccadic drift by altering the levels of innervation sent to the muscles during and after a saccade. The adaptive mechanism for suppression of post-saccadic drift is sensitive to optically-imposed post-saccadic retinal slip. Our previous work showed that a central neural mechanism attempted to compensate for this post-saccadic retinal slip by altering the gain and time constants of the neural components of saccadic innervation. This altered innervation led to a post-saccadic ocular drift that lessened the amount of post-saccadic retinal slip.

(c) Neither

We have previously shown that this central adaptive mechanism was dependent upon the cerebellum for its proper function. After a complete cerebellectomy, monkeys developed post-saccadic ocular drift, and were unable to compensate for post-saccadic retinal slip. The present work has now shown that the site of this functional dependency can be further localized to the cerebellar flocculi and paraflocculi.

After bilateral flocculectomy, monkeys developed post-saccadic ocular drift. When presented with optically-imposed post-saccadic retinal slip, the animals were only able to alter their saccadic innervations a little. These post-flocculectomy alterations were not large enough to compensate for the retinal slip. Furthermore, this ability was asymmetric, with reductions in gain being larger than increases.

PROJECT NUMBER DEPARTMENT OF HEALTH AND HUMAN SERVICES - PUBLIC HEALTH SERVICE NOTICE OF INTRAMURAL RESEARCH PROJECT Z01 EY 00045-08 LSR PERIOD COVERED October 1, 1985, to September 30, 1986 TITLE OF PROJECT (80 characters or less. Title must lit on one line between the borders.) Visuomotor Properties of Neurons in the Thalamus PRINCIPAL INVESTIGATOR (List other professional personnel below the Principal Investigator) (Name, title, laboratory, and institute affiliation) David Lee Robinson Ph.D. Research Physiologist LSR, NEI Others: John W. McClurkin Ph.D. Guest Worker LSR, NEI Jon Currie M.D. Neuro-Ophthalmologist LSR, NEI Mortimer Mishkin Ph.D. Laboratory Chief LNP, NIMH Marcie Golomb 0. D. Guest Worker LSR, NEI Richard Sherins M.D. Res. Endocrinologist RR, NICHD Edmond FitzGibbon M. D. Clinical Fellow LSR, NEI COOPERATING UNITS (# anv) LAB/BRANCH Laboratory of Sensorimotor Research Visuomotor Integration Section INSTITUTE AND LOCATION NEI, NIH, Bethesda, Maryland 20892 TOTAL MAN-YEARS PROFESSIONAL

SUMMARY OF WORK (Use standard unreduced type. Do not exceed the space provided.)

2.6

CHECK APPROPRIATE BOX(ES)

(a) Human subjects (a1) Minors (a2) Interviews

We have studied visual spatial attention in humans and monkeys to determine what brain areas are important and what each contributes to this cognitive function. Visual cues which precede visual targets can modulate reaction times to the targets. Cues on the same side as the target (valid cues) are associated with fast reaction times; cues on the opposite side (invalid cues) or weak illumination of the entire visual field (diffuse cues) are correlated with slow reaction times. The cues are hypothesized to control the direction of the subject's attention.

1.0

(b) Human tissues

OTHER

(c) Neither

1.6

Patients with damage to parietal cortex have slowed reaction times to targets which follow diffuse cues or to those cues which draw their attention to the intact visual field. These are both diffuse cues and one invalid cue. We have demonstrated the same deficit in the monkey after surgical removal of cortical area 7. This allows us to study directly a region of cortex in the monkey which is critical for visual spatial attention. Normal human controls or patients with temporal lobe damage do not have these problems. Humans with Alzheimer's dementia are extremely slow in all aspects of this task. Males with idiopathic hypogonadotropic hypogonadism have unusual patterns of responding on our attention task; they are slow in responding to all targets in their right visual field, independent of the preceding cue. They are also slow in responding to targets after diffuse cues.

These studies have helped to localize an area of the brain, the inferior parietal lobule, which is critical for visual spatial attention and helped to clarify its contribution to this cognitive process. In addition they have demonstrated a possible endocrine influence on visual behavior.

PROJECT NUMBER DEPARTMENT OF HEALTH AND HUMAN SERVICES - PUBLIC HEALTH SERVICE NOTICE OF INTRAMURAL RESEARCH PROJECT ZO1 EY 00109-06 LSR PERIOD COVERED October 1, 1985, to September 30, 1986 TITLE OF PROJECT (80 characters or less. Title must fit on one line between the borders.) Visual Motion Processing in the Primate Brain PRINCIPAL INVESTIGATOR (List other professional personnel below the Principal Investigator) (Name title laborator), and institute attitution PI: Robert H. Wurtz Ph.D. Chief LSR, NEI Others: Max R. Dursteler M.D. Visiting Scientist LSR, NEI Hidehiko Komatsu Ph.D. Visiting Scientist LSR, NEI Dwayne S. Yamasaki Ph.D. Guest Researcher LSR, NEI COOPERATING UNITS (if eny) LAB/BRANCH Laboratory of Sensorimotor Research Visuomotor Integration Section INSTITUTE AND LOCATION

OTHER

(c) Neither

2.0

SUMMARY OF WORK (Use standard unreduced type. Do not exceed the space provided.)

20892

2.8

(b) Human tissues

PROFESSIONAL

NEI, NIH, Bethesda, Maryland

TOTAL MAN-YEARS

CHECK APPROPRIATE BOX(ES)

☐ (a) Human subjects
☐ (a1) Minors
☐ (a2) Interviews

Visual motion processing in the cerebral cortex of the monkey is carried out in a series of specialized areas. We have previously investigated one of these areas, the middle temporal area (MT) and we have now explored the next area, the medial superior temporal area (MST). We investigated the relation of motion processing to the generation of an eye movement that is dependent on such motion processing, smooth pursuit eye movements. We found cells that discharge during pursuit eye movements fell into two groups. Those that were dependent upon visual stimulation lie in the foveal region of MT and lateral MST, while those related to the movement itself were in the lateral and dorsal MST. Punctate removal of cells in these areas of MST using a neurotoxin, ibotenic acid, showed a deficit in pursuit initiation for targets moving in the contralateral visual field (a retinotopic deficit) and a deficit in maintenance of pursuit as long as the target was moving toward the side of the lesion (a directional deficit). We interpret the visual response of cells in MT and the retinotopic deficit as being related to visual stimulation (retinal slip) and the pursuit related cells and the directional deficit as being related to an efference copy signal of pursuit eye movements.





	٠,	

	- 1
	- 1
	- 11
	- 11
	- 1
	- 1
	- 1
	- 1
	- 11
	- 11
	- 1
	- 1



http://nihlibrary.nih.gov

10 Center Drive Bethesda, MD 20892-1150 301-496-1080

